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FIELD OF THE INVENTION

The present invention relates to the design of trimeric polypeptides using polypeptide structural elements derived from the tetranectin protein family, and their use in rational de novo design and production of multi-functional molecules.

BACKGROUND OF THE INVENTION

Tetranectin is a Ca²⁺-binding trimeric C-type lectin which is present in blood plasma and from the extracellular matrix of certain tissues. The tetranectin group of proteins comprises tetranectin isolated from man and from mouse and the highly related C-type lectin homologues isolated from the cartilage of cattle (Neame and Boynton, database accession number PATCHX:u22298) and from reef shark (Neame et al., 1992, Neame et al., 1996 and database accession number p26258 and PIR2:A37289).

The mature tetranectin polypeptide chain of 181 amino acid residues is encoded in three exons as shown by molecular cloning and characterisation of the gene (Berglund & Petersen, 1992; Wewer & Albrechtsen, 1992). Exon 3 of the human tetranectin gene encodes a separate functional and structural unit, a single long-form so-called carbohydrate recognition domain (CRD), with three intra-chain disulphide bridges. The tetranectin CRD is considered to belong to a distinct class of C-type lectins (Day, 1994) clearly related to C-type lectins by sequence homology, conservation of disulphide topology (Fuhlendorff et al, 1987) and by the presence of an almost conserved suit of amino acid residues known to be involved in binding of calcium ions.

A published poster (Holtet et al 1996) has proposed tetranectin to be a trimer and that trimerisation is governed by the peptide encoded by exon 1. The peptide encoded by exon 1 was proposed to be "necessary and sufficient to govern trimerisa-

tion" whereas the polypeptide encoded by exon 2 was proposed as being "involved in lysine-sensitive binding to plasminogen".

SUMMARY OF THE INVENTION

5 It has surprisingly been found by the present inventors that the human tetranectin polypeptide (and derivatives thereof) is capable of forming very stable trimers which have a number of advantageous characteristics and uses. Notably, the tetranectin molecule includes a trimerising structural element which can be used as carrier of other chemical entities, thereby providing a carrier molecule of a hitherto unseen versatility.

Prior published knowledge in the field of providing trimerising polypeptides of choice includes the disclosure in WO 95/31540 by Hoppe and Reid which describes a trimerisation module derived from collectin coiled coil structures and its application in engineering of artificially trimerised proteins. Several interesting areas of application are common to that patent publication and to the present disclosure. However, in several ways the properties of the trimerisation modules derived from the tetranectin protein family as disclosed herein are markedly different in fundamental architecture and represent surprisingly improved properties in comparison with the collectin trimerisation unit:

25 (1) Although the spatial structures of both trimerisation modules at a superficial level appear as similar in that both are ternary coiled coil structures of roughly equivalent spatial size the structural basis for adopting this spatial configuration is markedly distinct between the two groups of proteins. In fact, it is so distinct that the common belief prior to the work of Holtet et al. on cross-linking of human tetranectin (Holtet et al., 1996) was that this family of proteins were tetrameric (hence the name). Accordingly, the sequences of the tetranectin family of trimerisation modules

does not conform to the declared common motif delineated for the collectin family (WO 95/31540, page 8).

- (2) The thermal stability of the tetranectin trimerisation module (as shown in the examples) is such that the trimer can be shown to exist even at about 60°C (Example 4, trimerised tetranectin) or at about 70°C (Example 3, trimerised ubiquitin), whereas a collectin trimer unit falls apart at about 50-55°C (WO 95/31540, Example 1, page 36 therein).
- (3) Whereas it remains uncertain whether the collectin trimerisation domain possibly allows attachment of fusion partners 10 at C-terminal ends of the trimerisation module, and whereas no example has been reported of successful or claimed successful attachment of a foreign protein (except for the GST fusion partner) to the N-terminal region of the collectin trimerisation module, the information disclosed herein demonstrates that the tetranectin trimerisation module is more versatile in that is allows attachment of foreign proteins to either, as well as to both, terminus or termini simultaneously (Examples 1-4). This has important consequences as the tetranectin trimerisation module may be deployed to construct 20 molecules that are able to interact (each end with a binding valency up to 3) simultaneously with two bulky interaction partners like e.g. cell surfaces.
- (4) The virtual absence of subunit exchange between monomers of a trimer that has been trimerised using the tetranectin trimerisation modules disclosed herein is by first principles of thermodynamics correlated with the surprisingly high thermal stability of the complex. It will hence be apparent that the advantages inherent to the "pick-and-mix" applications of the technology, as disclosed herein, may be used to much greater advantage because of the much longer shelf life expected for the heterofunctional products of the present invention.

The polypeptide constructs CIIH6FXTN123, H6FXTN123, H6FXTN12, and H6FXTN23 which all involve parts of the tetranectin molecule have previously been prepared (cf. e.g. WO 94/18227) but these constructs have all been provided with a view to facilitating expression and/or purification of the tetranectin derived moiety of the constructs. To the best of the inventors' knowledge no publications exist which reports any use of tetranectin derivatives as "building blocks" to which other chemical moieties advantageously could be coupled.

10 Hence, in its broadest aspect the present invention relates to a monomer polypeptide construct comprising at least one tetranectin trimerising structural element (hereinafter designated a TTSE) which is covalently linked to at least one heterologous moiety, said TTSE being capable of forming a stable complex with two other TTSEs, with the proviso that the heterologous moiety is different from any of the fusion proteins CIIH6FXTN123, H6FXTN123, H6FXTN12, H6FXTN23, the sequences of which are shown in SEQ ID NOs: 24-27. It is preferred that the heterologous moiety is one which does not exclusively facilitate expression and/or purification of the monomer polypeptide construct.

The invention further relates to oligomeric molecules comprised of at least two of such monomer polypeptide constructs, and the invention also relates to methods of preparing the monomer polypeptide constructs and the oligomers. The invention further relates to a kit comprising monomer polypeptide constructs in separate packages, ready for use in a "pick-and-mix" approach for use of the monomers. Finally, the invention also pertains to fragments which include nucleic acid sequences which encode the monomer polypeptide constructs, as well as to vectors and cells containing these nucleic acid fragments.

LEGENDS TO THE FIGURES

Fig. 1: Amino acid sequence of the amino terminal region of tetranectin.

Amino acid sequence (in one letter code) from El to L51 of tetranectin (SEQ ID NO: 7). Exon 1 comprises residues El to D16 and exon 2 residues V17 to V49, respectively. The alpha helix extends beyond L51 to K52 which is the C-terminal amino acid residue in the alpha helix.

Fig. 2: Alignment of the amino acid sequences of the trimerising structural element of the tetranectin protein family. Amino acid sequences (one letter code) corresponding to residue V17 to K52 comprising exon 2 and the first three residues of exon 3 of human tetranectin (SEQ ID NO: 7); murine tetranectin (Sørensen et al., Gene, 152: 243 -245, 1995); tetranectin homologous protein isolated from reef-15 shark cartilage (Neame and Boynton, 1992, 1996); and tetranectin homologous protein isolated from bovine cartilage (Neame and Boynton, database accession number PATCHX:u22298). Residues at a and d positions in the heptad repeats are listed in 20 boldface. The listed consensus sequence of the tetranectin protein family trimerising structural element comprise the residues present at a and d positions in the heptad repeats shown in the figure in addition to the other conserved residues of the region. "hy" denotes an aliphatic hydrophobic 25 residue.

Fig. 3: Construction of the expression plasmids pTH6FXtripa and pTH6FXtripb.

The amplified DNA fragments tripa and tripb harbouring the tetranectin amino acid sequence (SEQ ID NO: 7) from E1 to T48 and E1 to K52, respectively, fused in the 5' end to nucleotide sequences encoding a FX_a cleavage site IQGR (SEQ ID NO: 4) and the recognition sites for the restriction endonucleases BglII and KpnI, were cut with the restriction enzymes BclI and HindIII and ligated into the BamHI and

HindIII sites of the expression plasmid pT7H6 (Christensen et al., 1991) using standard procedures.

Fig. 4: Predicted amino acid sequence of the fusion proteins H6FXtripa (SEQ ID NO: 28) and H6FXtripb (SEQ ID NO: 29)
5 encoded by the expression plasmids pTH6FXtripa and pTH6FXtripb, respectively.

Fig. 5: Construction of the expression plasmids pTH6FXTN123 and pTCIIH6FXTN123.

The amplified DNA fragment corresponding to the full length,

mature tetranectin monomer (SEQ ID NO: 7) from E1 to V181

fused in the 5' end to nucleotide sequences encoding a FX_a

cleavage site IEGR (SEQ ID NO: 10) was cut with the restriction enzymes BamHI and HindIII and ligated into the corresponding sites of the expression plasmids pT7H6 (Christensen et al., 1991) and pTCIIH6 using standard procedures. pTCIIH6 was derived from pT7H6 by substitution of the NdeI - HindIII fragment of pT7H6 with the NdeI - HindIII fragment of pLcII (Nagai and Thøgersen, 1987) encoding the first 32 residues of the lambda cII protein MVRANKRNEALRIESALLNKIAMLGTEKTAEG (SEQ ID NO: 11) fused in the 3' end to a nucleotide sequence encoding the H6 sequence GSHHHHHHHGS(SEQ ID NO: 12).

Fig. 6: Predicted amino acid sequence of the fusion proteins H6FXTN123 (SEQ ID NO: 25) and CIIH6FXTN123 (SEQ ID NO: 24) encoded by the expression plasmids pTH6FXTN123 and pTCIIH6FXTN123, respectively.

Fig. 7: Construction of the expression plasmids pTH6FXTN12, pTH6FXTN23, and pTH6FXTN3.

The amplified DNA fragments corresponding to the tetranectin derivatives TN12 and TN3 from E1 to V49 and A45 to V181, respectively (SEQ ID NO: 7) fused in the 5' end to nucleotide sequences encoding the FX_a cleavage site IEGR (SEQ ID NO: 10) was cut with the restriction enzymes BamHI and HindIII and ligated into the corresponding sites of the expression plas-

mids pT7H6 (Christensen et al., 1991) using standard pro-

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cedures. The amplified DNA fragment corresponding to the tetranectin derivative TN23 from V17 to V181 (SEQ ID NO: 7) fused in the 5' end to nucleotide sequences encoding the FX_a cleavage site IQGR (SEQ ID NO: 4) was cut with the restriction enzymes BamHI and HindIII and ligated into the corresponding sites of the expression plasmids pT7H6 (Christensen et al., 1991) using standard procedures.

Fig. 8: Predicted amino acid sequence of the fusion proteins H6FXTN12 (SEQ ID NO: 26), H6FXTN23 (SEQ ID NO: 27), and H6FXTN3 (SEQ ID NO: 30) encoded by the expression plasmids pTH6FXTN12, pTH6FXTN12, respectively.

Fig. 9: Gel filtration analysis of TN123, TN23, and TN3
Analytical gel filtration of the recombinant tetranectin
derivatives TN123, TN23, and TN3 were performed on a Superose
15 12 HR 10/30 column (Pharmacia, Sweden) with a total volume of
25 ml in 100 mM NaCl and 50 mM Tris-HCl pH 8 and a flow rate
of 0.2 ml/min. Vertical bars at peak maxima identify elution
profiles for each of the three proteins.

Fig. 10: Cross-linking analysis of TN123 and CIIH6FXTN123.

20 Samples of TN123, CIIH6FXTN123 and mixtures of both were incubated with DMSI and analyzed by SDS-PAGE (12% gel).

Before addition of DMSI, protein mixtures were subjected to subunit exchange by incubation at 70°C for varying length of time. Protein marker of 94, 68, 43 and 30 kDa, top to bottom

25 (lane M). CIIH6FXTN123 fusion protein (lane 1). TN123 (lane 2). DMSI treated CIIH6FXTN123 (lanes 3 and 6). DMSI-treated TN123 (lane 4). Identical samples of DMSI treated mixtures of CIIH6FXTN123 and TN123 without heat exposure (lanes 5 and 7) and heat treated for 2.5 sec, 15 sec, 2.5 min. and 10 min., respectively, before treatment with DMSI (lanes 8-11).

Fig. 11: Cross-linking analysis of the recombinant tetranectin derivatives TN123, TN23, TN3, and H6FXTN12. The recombinant proteins TN123, TN23, TN3, H6FXTN12 or mixtures of TN123 and each of the other were analyzed by SDS-

PAGE. Protein marker of 94, 68, 43, 30, 20, and 14.4 kDa, top to bottom (lane M). TN123 cross-linked with DMSI (lane 1). TN123 and H6-rTN12 cross-linked with DMSI without and with heat treatment at 70°C for two min. (lanes 2 and 3). H6FXTN12 cross-linked with DMSI (lanes 4 and 5). Mixture of TN123 and H6FXTN12, no cross-linking (lane 6). Cross-linking of TN123 and TN23 without and with heat treatment at 70°C for two min. (lanes 7 and 8). Cross-linking of TN23 (lane 9). Mixture of TN123 and TN23 without cross-linking (lane 10). TN123 cross-linked by DMSI (lane 11). Cross-linking of TN123 and TN3 without and with heat treatment for two min. (lanes 12 and 13). Cross-linking of TN3 (Lane 14). Mixture of TN123 and TN3, no cross-linking (lane 15).

Fig. 12: Cross-linking based analysis of the trimer thermal stability.

In parallel experiments TN123 and the fusion protein
H6FXtripb-UB (SEQ ID NO: 31) were cross-linked with DMSI at
different temperatures and the samples analyzed by SDS-PAGE.
Protein marker of 94, 68, 43, 30, 20, and 14.4 kDa, top to

20 bottom (lane M). TN123 without cross-linking (lane 1). TN123
cross-linked with DMSI for 15 min. at 37°C, 50°C, 60°C, and
70°C (lanes 2 to 5), respectively. The fusion protein
H6FXtripb-UB (SEQ ID NO: 31) without cross-linking (lane 6).
H6FXtripb-UB cross-linked with DMSI for 15 min. at 37°C,

25 50°C, 60°C, and 70°C (lanes 7 to 10), respectively and
H6FXtripb-UB incubated at 70°C for 15 min. (lane 11).

Fig. 13: Construction of the expression plasmid pTH6FXtripb-UB.

The amplified DNA fragment comprising the nucleotide sequence (SEQ ID NO:16) encoding the ubiquitin amino acid sequence (SEQ ID NO: 19) from Q2 to G76 was cut with the restriction enzymes BclI and HindIII and ligated into the BamHI and HindIII sites of the expression plasmid pT7H6FXtripb (Example 1) using standard procedures.

Fig. 14: Predicted amino acid sequence of the fusion protein H6FXtripb-UB (SEQ ID NO: 31) encoded by the expression plasmid pTH6FXtripb-UB.

Fig. 15: Construction of the expression plasmid pTH6FXscFV (CEA6) tripb.

The DNA fragment, amplified with the primer pair SEQ ID NOs: 21 and 22, comprising the nucleotide sequence SEQ ID NO: 20 encoding the single chain antibody CEA6, scFV (CEA6), amino acid sequence from Q1 to A261 was cut with the restriction enzymes BamHI and KpnI and ligated into the BglII and KpnI sites of the expression plasmid pT7H6FXtripb (Example 1) using standard procedures.

Fig. 16: Predicted amino acid sequence of the fusion protein H6FXscFV(CEA6)tripb encoded by the expression plasmid pH6FXscFV(CEA6)tripb.

Fig. 17: Construction of the expression plasmid pTH6FXtripbscFX(CEA6).

The DNA fragment, amplified with the primer pairs having SEQ ID NO: 21 and 23, comprising the nucleotide sequence (SEQ ID NO: 20) encoding the single chain antibody CEA6, scFV (CEA6), amino acid sequence from Q1 to A261 was cut with the restriction enzymes BamHI and HindIII and ligated into the BamHI and HindIII sites of the expression plasmid pT7H6FXtripb (Example 1) using standard procedures.

- Fig. 18: Predicted amino acid sequence of the fusion protein H6FXtripbscFv(CEA6) encoded by the expression plasmid pH6FXtripbscFv(CEA6).
 - Fig. 19: Construction of the expression plasmid pTH6FXscFv(CEA6)tripbscFX(CEA6).
- The DNA fragment, amplified with the primer pair SEQ ID NO: 21 and 23, comprising the nucleotide sequence (SEQ ID NO: 20) encoding the single chain antibody CEA6, scFV (CEA6), amino acid sequence from Q1 to A261 was cut with the restriction

enzymes BamHI and HindIII and ligated into the BamHI and HindIII sites of the expression plasmid pT7H6FXscFv(CEA6)tripb (Example 4) using standard procedures.

Fig. 20: Predicted amino acid sequence of the fusion protein H6FXscFv(CEA6) tripbscFv(CEA6) (SEQ ID NO: 34) encoded by the expression plasmid pH6FXscFv(CEA6) tripbscFv(CEA6).

Fig. 21: Cross-linking analysis of the H6FXtripbscFv(CEA6) fusion protein (SEQ ID NO: 33). In parallel experiments the fusion proteins H6FXtripbscFv(CEA6) (SEO ID NO: 33) and TN123 were crosslinked at room temperature for 30 min. with 0 mg/ml, 0.5 mg/ml, 1.0 mg/ml, 1.5 mg/ml, and 2.0 mg/ml of DMSI, respectively. Lane 1: H6FXtripbscFv(CEA6) without DMSI, H6FXtripbscFv(CEA6) with 0.5 mg/ml DMSI (lane 2), 15 H6FXtripbscFv(CEA6) with 1.0 mg/ml DMSI (lane 3), H6FXtripbscFv(CEA6) with 1.5 mg/ml DMSI (lane 4) and H6FXtripbscFv(CEA6) with 2.0 mg/ml DMSI (lane 5). Protein marker of 94, 68, 43, 30, 20, and 14.4 kDa, top to bottom (lane M). Lane 6: TN123 without DMSI, TN123 with 0.5 mg/ml. DMSI (lane 7), TN123 with 1.0 mg/ml DMSI (lane 8) TN123 with 1.5 mg/ml DMSI (lane 9) and TN123 with 2.0 mg/ml DMSI (lane 10).

DETAILED DISCLOSURE OF THE INVENTION

The term "trimerising structural element" (TTSE) used in the present description and claims is intended to refer to the portion of a polypeptide molecule of the tetranectin family which is responsible for trimerisation between monomers of the tetranectin polypeptide. The term is also intended to embrace variants of a TTSE of a naturally occurring tetranectin family member, variants which have been modified in the amino acid sequence without adversely affecting, to any substantial degree, the trimerisation properties relative to those of the native tetranectin family member molecule. Specific examples of such variants will be described in

detail herein, but it is generally preferred that the TTSE is derived from human tetranectin, murine tetranectin, C-type lectin of bovine cartilage, or C-type lectin of shark cartilage. Especially preferred is monomer polypeptide constructs including at least one TTSE derived from human tetranectin.

The 49 residue polypeptide sequence encoded by exons 1 and 2 of tetranectin (Fig. 1) appears to be unique to the tetranectin group of proteins (Fig. 2) as no significant sequence homology to other known polypeptide sequences has been estab-10 lished. In preparation for experimental investigations of the architecture of tetranectin a collection of recombinant proteins was produced, the collection including complete tetranectin, the CRD domain (approximately corresponding to the polypeptide encoded by exon 3), a product corresponding to the polypeptide encoded by exons 2+3, a product correspon-15 ding to exons 1+2 (Holtet et al., 1996; Example 2). As detailed in Example 2 we now know differently: tetranectin is indeed a trimer, but the exon 2 encoded polypeptide is in fact capable of effecting trimerisation by itself as evi-20 denced by the observation that the recombinant protein corresponding to exons 2+3 is in fact trimeric in solution.

3D-structure analysis of crystals of full-length recombinant tetranectin (Nielsen et al., 1996; Nielsen, 1996; Larsen et al., 1996; Kastrup, 1996) has shown that the polypeptide encoded in exon 2 plus three residues encoded in exon 3 form a triple alpha helical coiled coil structure.

From the combination of sequence and structure data it becomes clear that trimerisation in tetranectin is in fact generated by a structural element (Fig. 2), comprising the amino acid residues encoded by exon two and the first three residues of exon 3 by an unusual heptad repeat sequence, that apparently is unique to tetranectin and other members of its group: This amino acid sequence (Fig. 2) is characterised by two copies of heptad repeats (abcdefg) with hydrophobic residues at a and d positions as are other alpha helical

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coiled coils. These two heptad repeats are in sequence followed by an unusual third copy of the heptad repeat, where glutamine 44 and glutamine 47 not only substitute the hydrophobic residues at both the a and d position, but are directly involved in the formation of the triple alpha helical coiled coil structure. These heptad repeats are additionally flanked by two half-repeats with hydrophobic residues at the d and a position, respectively.

The presence of beta-branched hydrophobic residues at a or d positions in alpha helical coiled coil are known to influence the state of oligomerisation. In the tetranectin structural element only one conserved valine (number 37) is present. At sequence position 29 in tetranectin no particular aliphatic residue appears to be preferred.

15 In summary, it is apparent that the triple stranded coiled coil structure in tetranectin to a large extent is governed by interactions that are unexpected in relation to those characteristic among the group of known coiled coil proteins.

The TTSEs form surprisingly stable trimeric molecules 20 (Examples 2, 3 and 4). The experimental observations, that (1) a substantial part of the recombinant proteins exists in the oligomeric state of - and can be cross-linked as trimeric molecules even at 70°C and (2) that exchange of monomers between different trimers can only be detected after 25 exposure to elevated temperature are evidence of a extremely high stability of the tetranectin trimerising structural element. This feature must be reflected in the amino acid sequence of the structural element. In particular, the presence and position of the glutamine containing repeat in the 30 sequential array of heptad repeats is, together with the presence and relative position of the other conserved residues in the consensus sequence (Fig. 2), considered important for the formation of these stable trimeric molecules. For most practical uses the cysteine residue 50 should be mutage-35 nized to serine, threonine, methionine or to any other amino

acid residue in order to avoid formation of an unwanted inter-chain disulphide bridge, which eventually would lead to uncontrolled multimerisation, aggregation and precipitation of a polypeptide product harbouring this sequence.

- In particular in conjunction with the trimer-stabilising exon 1 encoded polypeptide (tetranectin residues 1 to 16, see Example 2), the tetranectin trimerising structural element is a truly autonomous polypeptide module retaining its structural integrity and propensity to generate a highly stable 10 homotrimeric complex whether it is attached or not by a peptide bond at either or at both termini to other proteins. This unique property is demonstrated in the accompanying examples, which provide experimental proof, that polypeptide sequences derived from heterologous proteins may readily be trimerised when joined as fusion proteins to the tetranectin trimerising structural element. This remains valid irrespective of whether the heterologous polypeptide sequences are placed amino-terminally or carboxy-terminally to the trimerising element allowing for the formation of one molecular assembly containing up to six copies of one particular polypeptide sequence or functional entities, or the formation of one molecular assembly containing up to six different polypeptide sequences, each contributing their individual functional property.
- Since three TTSEs of naturally occurring human tetranectin forms up a triple alpha helical coiled coil, it is preferred that the stable complex formed by the TTSEs of the invention also forms a triple alpha helical coiled coil.
- The "tetranectin family" are polypeptides which share the

 consensus sequence shown in Fig. 2 or a sequence which are
 homologous at sequence level with this consensus sequence.
 Hence, monomer polypeptide constructs of the invention are
 preferred which comprise a polypeptide sequence which has at
 least 68% sequence identity with the consensus sequence shown
 in Fig. 2, but higher sequence identities are preferred, such

as at least 75%, at least 81%, at least 87%, and at least 92%.

By the term "heterologous moiety" is herein meant any chemical entity which can be linked covalently to a TTSE and to 5 which the TTSE is not natively covalently bound. Hence, the heterologous moiety can be any covalent partner moiety known in the art for providing desired binding, detection, or effector properties. The heterologous moiety can be a ligand binding structure such as a receptor molecule or the ligand 10 binding part of a receptor molecule, an antibody, an antigen binding antibody fragment, or a molecule having antibody characteristics such as e.g. the "diabodies" described in EP-A-0 672 142, or other ligand binding molecules such as avidin or streptavidin, or a lectin; a toxin such as ricin; a detectable label such as a fluorescence labelled molecule, a 15 radioactively labelled molecule, an enzymatically labelled molecule; an in situ activatable substance, such as a molecule which can be induced by radiation to be radioactively or chemically active; an enzyme such as a peroxidase; a 20 radioactive moiety such as a γ -, α -, β --, or β +-emitting molecule, e.g. a molecule comprising one or more radioactive isotopes selected from ¹⁴C, ³H, ³²P, ³³P, ²⁵S, ³⁸S, ³⁶Cl, ²²Na, $^{24}\mathrm{Na}$, $^{40}\mathrm{K}$, $^{42}\mathrm{K}$, $^{43}\mathrm{K}$, and any isotopes conventionally utilized for the purposes of facilitating detection of probes or the 25 purposes of providing localized radiation so as to effect cell death; a cytokine such as an interferon or a leukotriene; a non-proteinaceous polymer such as a polymeric alkaloid, 11.a polyalcohol, a polysaccharide, a lipid and a polyamine; a photo cross-linking moiety, i.e. a chemical 30 entity which effects cross-linking upon photo-activation; and a group facilitating conjugation of the monomer polypeptide construct to a target.

The heterologous moiety is preferably covalently linked to the TTSE by via a peptide bond to the N- or C-terminus of the 35 TTSE peptide chain, via a peptide bond to a side chain in the TTSE or via a bond to a cysteine residue, but any way of coupling covalently heterologous material to a polypeptide chain will be useful. The skilled person will know of such possibilities, e.g. by consulting the teachings of WO 95/31540 in this regard which are hereby incorporated by reference.

However, one interesting aspect of the invention relates to a monomer polypeptide construct of the invention comprising two heterologous moieties which are linked via peptide bonds to the N- and C-terminus, respectively. This approach introduces a number of possibilities in terms of e.g. linking larger entities with oligomers of the invention by having specific activities coupled to each end of the monomers (as explained in detail below, the oligomers of the invention may also utilise a version of this principle, where e.g. one N-terminus and one C-terminus of an oligomer are linked via peptide bonds to independent heterologous moieties).

In general, a complex between two or three monomers are described in the following way: three monomers having one TTSE each forms a trimer designated (1+1+1), whereas a dimer 20 formed between a monomer with two TTSEs and a monomer with one TTSE is designated (1+2). Other (undesired) trimers can of course be formed, e.g. (2+2+1), where two TTSEs are not "in use", but it is preferred that the oligomers of the invention use all of their available TTSEs during complex formation. It should also be noted that the term "monomer 25 polypeptide construct" is meant to designate a single polypeptide chain which may or may not have non-peptide groups coupled covalently to the polypeptide chain, whereas "dimeric polypeptide" or "dimer", "trimeric polypeptide" or "trimer" and "oligomer" (i.e. a dimer or trimer) in the 30 present context are meant to designate non-covalent complexes of monomer polypeptide constructs. I.e., the traditional definitions of monomers and multimers do not apply in the present specification and claims.

The TTSE as exemplified by exon 2 or exons 1 and 2 of human tetranectin, preferably so modified to allow only heterotrimerisation between dissimilar (1+1+1) or (1+2) (cf. the below discussion) subunits may be deployed as a general 5 affinity mediator, which can be coupled chemically to each member of a selection of target molecules. After such conjugation with TTSE the target molecules may be homo- or heterotrimerised as desired for any application. Similar deployment of dimerisation, using as one partner a polypeptide harbouring two TTSE sequences in-line, separated by a linker sequence of suitable length and character, may bet yet more advantageous, as in such case absolute control of stoichiometry in complex formation would be possible. Thus, an important embodiment of the invention is a monomer polypeptide construct of the invention comprising 2 TTSEs which are covalently linked by a spacer moiety which allows both of the 2 TTSEs to take part in complex formation with a third TTSE not being part of the monomer polypeptide construct, but equally important is the embodiment of the invention where the monomer polypeptide construct comprises one. single TTSE, so as to allow trimerisation between three monomers and hence providing the optimum degree of versatility with respect to the number of functional units which can be easily incorporated into one single complex.

In the embodiments where two TTSEs are present in the same monomer it is preferred that the spacer moiety has a length and a conformation which favours complex formation involving both of the two TTSEs which are covalently linked by the spacer moiety. In this way, problems arising from undesired formation of trimers of the formats (2+1+1), (2+2+2), and (2+2+1) (wherein only one TTSE of each monomer participates in complex formation) can be diminished. Design and preparation of suitable spacer moieties are known in the art and are conveniently effected by preparing fusion polypeptides having the format TTSE¹-Spacer-TTSE², where the spacer moiety is a polypeptide fragment (often a relatively inert one), so as to

avoid undesired reactions between the spacer and the surroundings or the TTSEs.

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One typical scenario, where such modification may be advantageous is the case of immunological detection where a chemical conjugate of an antibody with enzymes such as peroxidase is used for in situ staining purposes in tissue or on western blots.

A similar, but yet different, application example would be the deployment of TTSE to mediate conjugation of e.g. alkaline phosphatase and an oligonucleotide which would allow in situ identification of a given mRNA in a tissue sample concurrently with idenfification of any other mRNA molecule e.g. by interconnection of a second appropriate oligonucleotide and a signalling/visualisation molecule using e.g. the biotin - avidin/streptavidin affinity pair for conjugation. The point of having two or more selective affinity systems available for conjugating oligonucleotide probes and detector molecules is that as many different sequences may be detected simultaneously as there are affinity sets available.

In terms of chemistry required to exploit TTSE as a conjugating affinity-contributing agent, the peptide corresponding to exon 2 will have a sufficient affinity for most purposes, but incorporation of all, or some segment of the exon 1 polypeptide will serve to increase affinity and stability.

The properties of tetranectin mutants in which many of hydrophilic (e.g. lys and glu) residues that are largely exterior in the coiled coil structure have been replaced with alanine appear similar to the native protein, suggesting that is indeed possible without interfering very much with stability

of the trimeric structure to replace all glu, asp and lys residues by a combination of gln, asn, arg or ala, and thereby generate a sequence that, as an N-terminally blocked synthetic peptide, would be very easy to convert into a chemically stable active-ester component, e.g. an N-hydroxy

35 succinimide ester of an acetylated peptide, that could react

with (and thereby couple to) any exposed lysine side chain in a target molecule of interest. Such peptide synthesis, activation and coupling chemistry will be readily designed and applied by a person skilled in the art of peptide chemistry, 5 as will indeed any other conjugation chemistry, like the attachment and use of photo-activatable moieties like e.g. phenyl azides.

In conclusion, it seems that the most important structure in native TTSE is the consensus sequence shown in Fig. 2, and that large variations in the polypeptide chain may be 10 allowed. Hence, one advantageous embodiment of the monomer polypeptide construct of the invention is one where at least one amino acid residue selected from the group consisting of amino acid residue nos. 6, 21, 22, 24, 25, 27, 28, 31, 32, 35, 39, 41, 42, is/are substituted by any non-helix breaking 15 amino acid residue, the amino acid residue numbering referring to amino acid residues in SEQ ID NO: 7. All these residues have been shown not to be directly involved in the intermolecular interactions which stabilises the trimeric 20 complex between three TTSEs of native tetranectin monomers and it is therefore expected that these amino acids may be safely substituted with any amino acid which will not have an adverse effect on helix formation (notably proline, which introduces a rigid bend in a polypeptide chain).

Another advantageous embodiment of the monomer polypeptide construct of the invention is one which is free from any free amino and/or carboxy groups. This would favour synthesis of a TTSE by means of solid or liquid phase peptide synthesis, since there would be no need of introducing any protective groups during peptide synthesis.

Since the consensus sequence of Fig. 2 is believed important and since this consensus sequence embraces the above-discussed heptad repeat, it is according to the invention preferred that the TTSE comprises a repeated heptad having the formula a-b-c-d-e-f-g (N to C), wherein residues a and d

generally are hydrophobic amino acids. However, since "a" and "d" in the third of the complete heptads of all known members of the tetranectin family are constituted of glutamine, it is most preferred that the TTSE comprises the heptad repeated 3 times and that the last occurrence of the heptad has a glutamine residue corresponding to residues a and d.

Since exon 2 of native members of the tetranectin family seems to contain the necessary elements to effect stable trimerisation, it is preferred that the monomer polypeptide construct is free of substantial parts of tetranectin which is encoded by exon 3 and/or lacks substantial parts of tetranectin which is encoded by exon 1. However, since exon 1 encoded material seems to stabilise the trimeric native tetranectin, it is especially preferred that all or part of exon 1 is part of the monomer polypeptide construct, and it also seems to be rational to include the first three amino acids encoded by exon 3, since these are known to take part of the formation of the native triple alpha helical coiled coil in human tetranectin.

One particularly interesting embodiment of the invention is the possibility of designing oriented molecular assemblies, where one or more functional entities are located N-terminally to the trimerising element and one or more functional entities are located C-terminally to the element. Such types of design may be particularly advantageous where a certain relative ratio is desired among the different functional entities included in a specific molecular unit. Such type of design may in addition be used if one or more functional entities for either structural or functional reasons appear incompatible within the same construct. Such may be the case if one or more of the functional entities are expressed by large or bulky protein domains which for steric reasons might prevent formation of the trimeric molecular unit due to sterical constraints.

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The possibility of constructing bi-polar three-way fusion proteins in which one functionality is placed N-terminally to the coiled coil structure and a different functionality is placed C-terminally is additionally advantageous in applications where large spatial separation between the two functionalities are desirable for optimal function. Examples of such application are e.g. the deployment of binding domains (e.g. antibody-derived binding modules) for recognition and binding to binding sites located at or close to large structures like cell membranes in cases where it is advantageous to allow for binding of the other end of the trimerised molecule to a different, but also bulky target.

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Hence, as discussed above, the oligomers of the invention may be used to join e.g. bulky surfaces by the oligomer according to the invention comprising at least one heterologous moiety which is positioned N-terminally to a TTSE and at least one heterologous moiety which is positioned C-terminally to a TTSE. The two heterologous moieties can be either part of the same monomer polypeptide construct or parts of two separate monomer polypeptide constructs.

The extraordinarily high stability of any trimer containing the tetranectin trimerisation module under physiological buffer and temperature condition (i.e. absence of denaturant, temperature not exceeding 40°C) in combination with the ease by which exchange of monomer subunits between trimers can be effected by incubation at moderately elevated temperature or in the presence of denaturants provide for a unique opportunity to deploy the trimerisation module as a vehicle to allow the construction of "pick-and-mix" conjugates prepared from previously fabricated collections of homotrimeric molecules. To illustrate the versatility of this design opportunity by way of theoretical example, let us assume that (1) a collection of twenty different antibody constructs (e.g. in the format of single-chain Fv) each of its own characteristic binding specificity, has been selected and then turned into homo-trimeric molecules by fusion to a tetranectin trimerisa-

tion module, and let us also assume that a set of twenty different effector molecules (e.g. toxin domains) have similarly been prepared and also conjugated to the tetranectin trimerisation module. A user provided with prefabricated collections of twenty different antibody constructs and twenty different toxin constructs - 40 different reagents in all - has the opportunity then to prepare 400 different toxin-antibody conjugates, simply by mixing a first preferred component from one reagent collection with a second preferred 10 reagent from the other collection and then subject this binary mixture to conditions, i.e. gentle heating or incubation with a suitable level of denaturant, to accomplish subunit exchange among all trimeric molecular species in the mixture. After the subunit exchange step the desired heterobifunctional reagent will be present in the mixture as a 15 major component of the mixture and may then be deployed as such to accomplish a given purpose or, alternatively apply a simple purification step to isolate his favoured heterofunctional binary reagent from any remaining mono-functional trimer species by a simple standard protein purification 20 step, easily designed using standard techniques known in the field of protein purification.

A further enhancement of the versatility of the "pick-andmix" technology may be achieved by including a specific 25 affinity purification tag on each array of trimerisation module - probe/effector/indicator conjugate, fused directly in-line or, alternatively, fused via a cleavable linker (a polypeptide segment containing e.g. a factor $\mathbf{X}_{\mathbf{a}}$ or an enterokinase recognition/cleavage site) to the affinity tag. 30 More specifically, if each of three libraries were tagged with affinity handles a, b and c, respectively, that were recognised by binding substances A, B and C, respectively, pure heterotrimers, composed of one member of each library, could be obtained in a three-step affinity purification procedure designed to allow selective recovery of only such trimers that exhibit affinity for substances A and B and C. If, for any reason, subsequent removal of the affinity tags

were desirable, and the constructs had been prepared to include cleavable linkers, isolation of the pure heterotrimer, liberated from all affinity tags, could be accomplished by three further affinity purification steps, arranged to isolate only material that would bind to neither substance A nor substance B nor substance C.

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Obviously, the scope of "pick-and-mix" design of user-preparable heterofunctional complexes apply not only to the formation of binary hetero-functionality, but would apply by logic extension to the formation of ternary hetero-functionality:

To envisage the wealth of possibilities that are inherent to the concept of ternary hetero-functionality in a further theoretical example along the lines given above, three sets of reagent collections, each comprising 20 different functional characteristics, i.e. a collection of in toto 60 different homotrimers would allow "pick-and-mix" preparation of 8,000 different tri-functional molecules.

The basic tetranectin trimerisation module will, essentially indiscriminately, form homo- and hetero-trimers with any 20 molecule that also contains a trimerisation module. For some applications it may be advantageous to have available specially engineered derivatives of the tetranectin trimerisation module, which have been reengineered to disallow homotrimer formation and hence only allow hetero-trimerisation. Thus, an important embodiment of the monomer polypeptide construct of the invention is constructed/reengineered so as to disfavour formation of complexes between identical TTSEs; this also has the implication that oligomers of the invention can advantageously be comprised of monomer polypeptide constructs which are designed so as to disfavour formation of trimers including two monomer polypeptide constructs having identical TTSEs.

The design/reengineering may be accomplished by introduction of amino acid substitution at sites in the polypeptide inti35 mately involved in the formation and stability of the trimer

and, simultaneously, in a different construct introduce a compensatory amino acid substitution, all in all removing symmetry between individual monomer components of the triple helical structure so that the structural complementarity profile only allows the formation of hetero-trimers, but is incompatible with some or each of the homotrimer species.

A yet different way to deploy the tetranectin trimerisation module as a vehicle to accomplish rational formation of bifunctionalisation would require the interconnection of the C-terminus of one monomer to the N-terminus of a second 10 monomer in the triple-helical structure. The basic requirement for such an intervening polypeptide is, that allowed spatial distances between its N- and C-termini must be compatible with the spacing inherent to the structural requirements given by the architecture of the tetranectin trimerisation module. The construction of an intervening connecting polypeptide allowed according to such criteria would be readily accomplished by an average person skilled in the art of protein engineering, as an ample collection of examples of the deployment of, usually flexible, spacer sequences are 20 known both in nature and in designed proteins. Due to the expected entropic contribution to interaction energy in a molecule in which two of the three tetranectin trimerisation module components are covalently tied together, such a mo-25 lecule would show great preference for selecting any molecule containing only a single copy of the tetranectin trimerisation module component, as this selection would be energetically favoured. Hence, conjugation of one functional protein component to a suitably selected covalently dimerised tetranectin trimerisation module component and conjugation of a 30 different functional protein component to a single-copy element of the trimerisation sequence would provide for the preferential formation of a 1:1 bifunctional complex and suppression of formation of any other complex.

35 The monomers of the invention may be prepared by methods generally known in the art, using exclusively or in combina-

tion the techniques of recombinant protein production, peptide synthesis (liquid phase or solid phase), and traditional chemical coupling of heterologous moieties to a peptide chain or to specific residues therein. Hence the 5 invention also relates to a method of preparing the monomer polypeptide construct of the invention, the method comprising

- isolating the monomer polypeptide construct from a culture comprising a host cell which carries and expresses a nucleic acid fragment which encodes the monomer polypeptide construct,
- synthesizing, by means of chemical peptide synthesis, the monomer polypeptide construct and subsequently isolating the monomer polypeptide construct from the reaction mixture,
- 15 preparing a TTSE in a culture comprising a host cell which carries and expresses a nucleic acid fragment which encodes the TTSE, subsequently linking covalently at least one heterologous moiety to the TTSE, and thereafter isolating the resulting monomer polypeptide construct, or
- 20 synthesizing, by means of chemical peptide synthesis, a TTSE, subsequently linking covalently at least one heterologous moiety to the TTSE, and thereafter the isolating the resulting monomer polypeptide construct from the reaction mixture.
- 25 and optionally subjecting the monomer polypeptide construct to further processing.

The nucleic acid fragment which is mentioned above is also a part of the invention and is defined as a nucleic acid fragment in isolated form which encodes a TTSE as defined herein 30 or which encodes the polypeptide part of a monomer polypeptide construct according to the invention, with the proviso that the nucleic acid fragment is different from one

that encodes native members of the tetranectin family, and that the nucleic acid fragment is different from one that encodes any of the fusion proteins CIIH6FXTN123, H6FXTN123, H6FXTN123, the sequences of which are shown in SEQ ID NOs: 24-27.

The above mentioned host cell (which is also a part of the invention) can be prepared by traditional genetic engineering techniques which comprises inserting a nucleic acid fragment (normally a DNA fragment) encoding the polypeptide part of a monomer polypeptide construct of the invention into a suit-10 able expression vector, transforming a suitable host cell with the vector, and culturing the host cell under conditions allowing expression of the polypeptide part of the monomer polypeptide construct. The nucleic acid fragment encoding the polypeptide may be placed under the control of a suitable 15 promoter which may be inducible or a constitutive promoter. Depending on the expression system, the polypeptide may be recovered from the extracellular phase, the periplasm or from the cytoplasm of the host cell.

Suitable vector systems and host cells are well-known in the art as evidenced by the vast amount of literature and materials available to the skilled person. Since the present invention also relates to the use of the nucleic acid fragments of the invention in the construction of vectors and in host cells, the following provides a general discussion relating to such use and the particular considerations in practising this aspect of the invention.

In general, of course, prokaryotes are preferred for the initial cloning of nucleic sequences of the invention and constructing the vectors useful in the invention. For example, in addition to the particular strains mentioned in the more specific disclosure below, one may mention by way of example, strains such as *E. coli* K12 strain 294 (ATCC No. 31446), *E. coli* B, and *E. coli* X 1776 (ATCC No. 31537). These

examples are, of course, intended to be illustrative rather than limiting.

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Prokaryotes are also preferred for expression, since efficient purification and protein refolding strategies are available. The aforementioned strains, as well as *E. coli* W3110 (F-, lambda-, prototrophic, ATCC No. 273325), bacilli such as *Bacillus subtilis*, or other enterobacteriaceae such as *Salmonella typhimurium* or *Serratia marcesans*, and various *Pseudomonas* species may be used.

In general, plasmid vectors containing replicon and control sequences which are derived from species compatible with the host cell are used in connection with these hosts. The vector ordinarily carries a replication site, as well as marking sequences which are capable of providing phenotypic selection in transformed cells. For example, *E. coli* is typically transformed using pBR322, a plasmid derived from an *E. coli* species (see, e.g., Bolivar et al., 1977). The pBR322 plasmid contains genes for ampicillin and tetracycline resistance and thus provides easy means for identifying transformed cells.

The pBR plasmid, or other microbial plasmid or phage must also contain, or be modified to contain, promoters which can be used by the microorganism for expression.

Those promoters most commonly used in recombinant DNA construction include the B-lactamase (penicillinase) and lactose promoter systems (Chang et al., 1978; Itakura et al., 1977; Goeddel et al., 1979) and a tryptophan (trp) promoter system (Goeddel et al., 1979; EPO Appl. Publ. No. 0036776). While these are the most commonly used, other microbial promoters have been discovered and utilized, and details concerning their nucleotide sequences have been published, enabling a skilled worker to ligate them functionally with plasmid vectors (Siebwenlist et al., 1980). Certain genes from prokaryotes may be expressed efficiently in E. coli from their own promoter sequences, precluding the need for addition of another promoter by artificial means.

In addition to prokaryotes, eukaryotic microbes, such as yeast cultures may also be used. Saccharomyces cerevisiase, or common baker's yeast is the most commonly used among eukaryotic microorganisms, although a number of other strains are commonly available. For expression in Saccharomyces, the plasmid YRp7, for example, is commonly used (Stinchcomb et al., 1979; Kingsman et al., 1979; Tschemper et al., 1980). This plasmid already contains the trpl gene which provides a selection marker for a mutant strain of yeast lacking the ability to grow in tryptophan for example ATCC No. 44076 or PEP4-1 (Jones, 1977). The presence of the trpl lesion as a characteristic of the yeast host cell genome then provides an effective environment for detecting transformation by growth in the absence of tryptophan.

Suitable promoting sequences in yeast vectors include the promoters for 3-phosphoglycerate kinase (Hitzman et al., 1980) or other glycolytic enzymes (Hess et al., 1968; Holland et al., 1978), such as enolase, glyceraldehyde-3-phosphate dehydrogenase, hexokinase, pyruvate decarboxylase, phosphofurate mutase, glucose-6-phosphate isomerase, 3-phosphoglycerate mutase, pyruvate kinase, triosephosphate isomerase, phosphoglucose isomerase, and glucokinase. In constructing suitable expression plasmids, the termination sequences associated with these genes are also ligated into the expression vector 3' of the sequence desired to be expressed to provide polyadenylation of the mRNA and termination.

Other promoters, which have the additional advantage of transcription controlled by growth conditions are the promoter region for alcohol dehydrogenase 2, isocytochrome C, acid phosphatase, degradative enzymes associated with nitrogen metabolism, and the aforementioned glyceraldehyde-3-phosphate dehydrogenase, and enzymes responsible for maltose and galactose utilization. Any plasmid vector containing a yeast-compatible promoter, origin of replication and termination sequences is suitable.

In addition to microorganisms, cultures of cells derived from multicellular organisms may also be used as hosts. In principle, any such cell culture is workable, whether from vertebrate or invertebrate culture. However, interest has been 5 greatest in vertebrate cells, and propagation of vertebrate in culture (tissue culture) has become a routine procedure in recent years (Tissue Culture, 1973). Examples of such useful host cell lines are VERO and HeLa cells, Chinese hamster ovary (CHO) cell lines, and W138, BHK, COS-7 293 and MDCK cell lines.

Expression vectors for such cells ordinarily include (if necessary) an origin of replication, a promoter located in front of the gene to be expressed, along with any necessary ribosome binding sites, RNA splice sites, polyadenylation site, and transcriptional terminator sequences.

For use in mammalian cells, the control functions on the expression vectors are often provided by viral material. For example, commonly used promoters are derived from polyoma, ... Adenovirus 2, and most frequently Simian Virus 40 (SV40). The early and late promoters of SV40 virus are particularly useful because both are obtained easily from the virus as a fragment which also contains the SV40 viral origin of replication (Fiers et al., 1978). Smaller or larger SV40 fragments may also be used, provided there is included the approximately 250 bp sequence extending from the HindIII site toward the BglI site located in the viral origin of replication. Further, it is also possible, and often desirable, to utilize promoter or control sequences normally associated with the desired gene sequence, provided such control sequences are compatible with the host cell systems.

An origin of replication may be provided either by construction of the vector to include an exogenous origin, such as may be derived from SV40 or other viral (e.g., Polyoma, Adeno, VSV, BPV) or may be provided by the host cell chromosomal replication mechanism. If the vector is inte-

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grated into the host cell chromosome, the latter is often sufficient.

Upon production of the polypeptide monomer constructs it may be necessary to process the polypeptides further, e.g. by introducing non-proteinaceous functions in the polypeptide, by subjecting the material to suitable refolding conditions (e.g. by using the generally applicable strategies suggested in WO 94/18227), or by cleaving off undesired peptide moieties of the monomer (e.g. expression enhancing peptide fragments which are undesired in the end product).

In the light of the above discussion, the methods for recombinantly producing the monomer polypeptide construct of the invention are also a part of the invention, as are the vectors carrying and/or being capable of replicating the nucleic acids according to the invention in a host cell or a cell-line. According to the invention the expression vector can be e.g. a plasmid, a cosmid, a minichromosome, or a phage. Especially interesting are vectors which are integrated in the host cell/cell line genome after introduction in the host.

Another part of the invention are transformed cells (useful in the above-described methods) carrying and capable of replicating the nucleic acid fragments of the invention; the host cell can be a microorganism such as a bacterium, a yeast, or a protozoan, or a cell derived from a multicellular organism such as a fungus, an insect cell, a plant cell, or a mammalian cell. Especially interesting are cells from the bacterial species *Escherichia*, *Bacillus* and *Salmonella*, and a preferred bacterium is *E. coli*.

Yet another part of the invention relates to a stable cell line producing the polypeptide part of a monomer polypeptide construct according to the invention, and preferably the cell line carries and expresses a nucleic acid of the invention.

On the basis of the above discussions it will be clear to the skilled person that also the oligomers resulting from the complex formation between the monomer constructs of the invention are important parts of the invention. Hence the invention also pertains to an oligomer which is comprised of two monomer polypeptide constructs according to the invention which comprises at least three TTSEs, or which is comprised of three monomer polypeptide constructs according to the invention which each only contain one single TTSE.

As is explained herein and shown in the examples, the oligomers of the invention are stable at temperatures up to 70°C and it is therefore especially preferred that the oligomers of the invention are stable at temperatures above physiological ones, e.g. that the oligomers are stable in the temperature range 50-70°C.

Also a part of the invention is a method for preparing a dimeric oligomer of the invention which comprises

- admixing a monomer polypeptide construct which includes two TTSEs (construct 1) with a monomer polypeptide construct which includes only one TTSE (construct 2),
- effecting the two TTSE's of construct 1 to complex with the TTSE of construct 2 (this can be done by thermal treatment, i.e. heating to a temperature which ensures denaturation followed by subsequent cooling allowing renaturation, or this can be done by denaturing/renaturing effected by changes in the chemical environment), and
- isolating the resulting dimer and optionally subjecting the dimer to further processing (cf. the above discussion of further processing, but it should also be mentioned that the further processing could include non-covalent coupling of interesting and relevant moieties to the dimeric oligomer).

Consequently, the method for producing a trimeric oligomer is also a part of the invention and comprises the steps of

- admixing three monomer polypeptide constructs of the invention with each other,
- 5 effecting complex formation between one TTSE of each monomer polypeptide construct, and
 - isolating the resulting trimer and optionally subjecting the trimeric oligomer to further processing.

The considerations applying to complex formation and further processing mentioned above apply to this method also.

Finally, in view of the detailed discussion above of the "pick-and-mix" aspect of the invention, the invention also pertains to a kit comprising

- a first package comprising at least one container means,
 each at least one container means containing a monomer
 polypeptide construct of the invention,
 - a second package comprising at least one container means, each at least one container means in the second package containing a monomer polypeptide construct of the invention, the second package being different from the first package with respect to choice and/or number of monomer polypeptide constructs included therein, and optionally
- a third package comprising at least one container means, each at least one container means in the third package containing a monomer polypeptide construct of the invention, the second package being different from the first and second packages with respect to choice and/or number of monomer polypeptide constructs included therein.

It is preferred that the at least one container means in each package contains mutually distinct monomer polypeptide constructs, and it is especially preferred that all container means comprised in the kit comprises mutually distinct polypeptide constructs.

EXAMPLE 1

Design and construction of the pTH6trip E. coli expression vectors for the production of trimerised chimeric fusion proteins.

10 The plasmid clone pT7H6FXTN123 (Example 2) was used as template for amplification in two Polymerase Chain Reactions (PCR) (Saiki et al., 1988) with the primer pairs trip-N (SEQ ID NO: 1) and trip-Ca (SEQ ID NO: 2) and trip-N (SEQ ID NO: 1) and trip-Cb (SEQ ID NO: 3), respectively. The amplified 15 DNA fragments, tripa, comprising nucleotide sequences encoding an IQGR cleavage site for the restriction protease FX : (SEQ ID NO: 4) followed by two sites for the restriction nucleases BglII and KpnI, the nucleotide sequence encoding the tetranectin polypeptide sequence for Glu 1 to Lys 52 (SEQ 20 ID NO: 5) followed by recognition sites for the three restriction endonucleases BamHI, HindIII, and EcoRI, respectively, and tripb, comprising nucleotide sequences encoding an IQGR cleavage site for the restriction protease FXa (SEQ ID NO: 4) followed by two sites for the restriction nucleases 25 BglII and KpnI, the nucleotide sequence encoding the tetranectin polypeptide sequence for Glu 1 to Val 49 (SEQ ID NO: 6) followed by recognition sites for the three restriction endonucleases BamHI, HindIII, and EcoRI, respectively, were subcloned into the plasmid pT7H6 (Christensen et al., 1991), yielding pTtripa and pTtripb, respectively (Figs. 3 and 4).

EXAMPLE 2

Tetranectin, localisation of the trimerising structural element and stability of the triple alpha helical coiled coil.

The cDNA encoding the reading frame corresponding to the mature tetranectin single chain (SEQ ID NO: 7) was cloned by specific amplification in a Polymerase Chain Reaction (PCR) (Saiki et al., 1988) of the nucleotide sequences from amino acid residue Glu1 to Val181 using 1st strand oligo-dT primed cDNA synthesized from total human placental RNA as template. Primers used in the PCR were SEQ ID NO: 8 and SEQ ID NO: 9. RNA extraction and cDNA synthesis were performed using standard procedures. The amplified reading frame encoding the monomer subunit of tetranectin was at the 5'-end, via the PCR-reaction, linked to nucleotide sequences encoding the 15 amino acid sequence SEQ ID NO: 10 which constitute an IEGR cleavage site for the bovine restriction protease FX, (Nagai, and Thøgersen, 1987). A glycine residue was, due to the specific design of the 5'-PCR primer (SEQ. ID NO. 8), 20 inserted between the C-terminal arginine residue of the FX, cleavage site (SEQ ID NO. 10) and the tetranectin Glu1-residue. The amplified DNA fragment was subcloned into the E. coli expression vector pT7H6 (Christensen et al., 1991) producing the plasmid pT₇H₆FX-TN123 expressing the tetranectin monomer H6FXTN123 (SEQ ID NO: 25) and into pT7CIIH6, 25 which is a derivative of pT_7H_6 , where the amino-terminal 32 amino acid residues of the lambda CII protein (SEQ ID NO. 11) are inserted 5' of the six histidine residues (SEQ ID NO. 12) as outlined in Fig. 5, yielding pT₇CIIH₆FX-TN123 expressing the tetranectin fusion protein CIIH6FXTN123 (SEQ ID NO: 24). The amino acid sequence of the expressed proteins are shown in Fig. 6 (in SEQ ID NO: 7 is given the amino acid sequence of the mature tetranectin protein). Furthermore three additional derivatives of tetranectin were constructed (Fig. 8): H6FXTN12 comprising the tetranectin amino acid residues Glu1 to Val49 (SEQ ID NO: 6), H6FXTN23 comprising the tetranectin

amino acid residues Val17 to Val181 (SEQ ID NO: 7), and H6FXTN3 (SEQ ID NO: 30) comprising the tetranectin amino acid residues Ala45 to Val181 (SEQ ID NO: 7). These three tetranectin derivatives were constructed by specific amplification 5 in a PCR using pT₇H₆FX-TN123 as template and the primer-pairs SEQ ID NO: 8 and SEQ ID NO: 13, SEQ ID NO: 14 and SEQ ID NO: 9, and SEQ ID NO: 15 and SEQ ID NO: 9, respectively. The amplified DNA fragments were subcloned into the E. coli expression vector pT_7H_6 producing the plasmids $pT_7H_6FX-TN12$, $pT_7H_6FX-TN23$, and $pT_7H_6FX-TN3$, respectively (Fig. 7).

To prepare recombinant tetranectin and its derivatives, each of the plasmids pT₇H₆FX-TN123, pT₇CIIH₆FX-TN123, pT₇H₆FX-TN12, $pT_7H_6FX-TN23$, and $pT_7H_6FX-TN3$ were grown in medium scale (4 x 1 litre; 2xTY medium, 5 mM MgSO₄ and 100 μ g ampicillin) in E.coli BL21 cells, as described by Studier et al. (1990). 15 Exponentially growing cultures at 37°C were at OD₆₀₀ 0.8 infected with bacteriophage lambda CE6 at a multiplicity of approximately 5. Cultures were grown at 37°C for another three hours and the cells harvested by centrifugation.

- 20 Cells were resuspended in 150 ml of 0.5 M NaCl, 10 mM Tris-HCl pH 8, and 1 mM EDTA pH 8. Phenol (100 ml adjusted to pH 8) was added and the mixture sonicated to extract the total protein. Protein was precipitated from the phenol phase by 2.5 volumes of ethanol and centrifugation.
- 25 The protein pellet was dissolved in a buffer containing 6M guanidinium chloride, 50 mM Tris-HCl pH 8 and 0.1 M dithioerythriol. Following gel filtration on Sephadex G-25 (Pharmacia, Sweden) into 8 M Urea, 1 M NaCl, 50 mM Tris-HCl pH 8 and 10 mM 2-mercaptoethanol, the crude protein preparation was applied to a Ni²⁺ activated NTA-agarose column (Ni²⁺NTA-agarose, 75 ml pre-washed with 8 M urea, 1 M NaCl, 50 mM Tris-HCl pH 8, and 10 mM 2-mercaptoethanol) for purification (Hochuli et al., 1988) and refolding of the fusion proteins, H6FXTN123, CIIH6FXTN123, H6FXTN12, H6FXTN23, and H6FXTN3.

For this study we chose to prepare our own Ni²⁺NTA-agarose matrix. A carbodiimide coupling of the N-(5-amino-1-carboxy-pentyl) iminodiacetic acid metal ligand (synthesis route as described by Döbeli & Hochuli (EP-A-0 253 303)) to a rigid agarose matrix (Sepharose CL-6B, Pharmacia, Sweden) was performed:

8 g of N-(5-amino-1-carboxypentyl)iminodiacetic acid from the synthesis procedure in 50 ml was adjusted to pH 10 by addition of 29 g of Na₂CO₃(10 H₂O) and added to a stirred suspension of activated Sepharose CL-6B in 1 M Na₂CO₃. Reaction was allowed overnight. The Sepharose CL-6B (initially 100 ml suspension) was activated after removal of water by acetone with 7 g of 1,1'-carbonyldiimidazol under stirring for 15 to 30 min. Upon activation the Sepharose CL-6B was washed with acetone followed by water and 1 M Na₂CO₃.

The NTA-agarose matrix was loaded into a column and "charged" with Ni²⁺ by slowly passing through 5 column volumes of a 10% NiSO₄ solution. The amount of Ni²⁺ on the NTA-agarose matrix, prepared by this procedure, has been determined to 14 μmol per ml matrix. After charging the Ni²⁺NTA-agarose column was washed with two column volumes of water, one column volume of 1 M Tris-HCl pH 8 and two column volumes of loading buffer before stirred mixing of the Ni²⁺NTA-agarose matrix with the crude protein extracts in a breaker for 15 to 30 min. All buffers prepared for liquid chromatography were degassed under vacuum prior to addition of reductant and/or use.

The Ni²⁺NTA-agarose matrix - crude extract mixture was packed in standard glass columns for liquid chromatography (internal diameter: 2.6 cm) to a volume of approximately 40 ml. The columns were washed with 200 ml of 8 M urea, 1 M NaCl, 50 mM Tris-HCl pH 8, and 10 mM 2-mercaptoethanol (Buffer I) and 100 ml 6 M guanidinium chloride, 50 mM Tris-HCl pH 8 and 10 mM 2-mercaptoethanol (Buffer II) and the adsorbed tetranectin derived fusion proteins H6FXTN123, H6CIIFXTN123, H6FXTN23,

and H6FXTN3 refolded using the cyclic refolding procedure as described (Thøgersen et al., WO 94/18227).

The fusion protein H6FXTN12 was refolded by removing the guanidinium chloride and 2-mercaptoethanol of buffer II in a gradient over 5 column volumes into 50 mM Tris-HCl pH 8 and 0.5 M NaCl. After completion of the refolding procedures the tetranectin derived fusion proteins were eluted from the Ni²⁺NTA-agarose columns with a buffer containing 0.5 M NaCl, 50 mM Tris-HCl, 25 mM EDTA pH 8. The tetranectin fusion proteins H6FXTN123, H6FXTN23, and H6FXTN3 were cleaved with 10 FX_a at 4°C overnight in a molar ratio of 1:300. After FX_a cleavage the protein samples were concentrated 10 fold by ultrafiltration on YM10 membranes (Amicon). After ten times dilution of the protein sample with 2 mM CaCl2, the recombi-15 nant tetranectin derivatives TN123, TN23, and TN3 were isolated by ion-exchange chromatography on Q-Sepharose (Pharmacia, Sweden) in a linear gradient over 10 column volumes from 10 mM Tris-HCl pH 8, 2 mM CaCl2 to 10 mM Tris-HCl pH 8, 2 mM CaCl₂, and 0.5 M NaCl. After elution from the Ni²⁺NTA-agarose 20 columns the fusion proteins H6CIIFXTN123 and H6FXTN12 were likewise concentrated 10 fold by ultrafiltration on YM10 membranes and gelfiltrated into buffer containing 25 mM Tris-HCl pH 8, 25 mM NaCl, and 2 mM CaCl2, before purification of correctly folded monomer by ion-exchange chromatography on 25 Q-Sepharose as described.

Recombinant full length tetranectin (TN123) produced by these procedures have been analyzed with respect to binding to plasminogen kringle 4 and immobilised fucoidan, expression of antigenic sites, and localization of disulphide bridges. In all criteria tested the produced TN123 behaved identically to isolated naturally human tetranectin (data not shown). Furthermore TN123 and TN3 have been crystallized (Kastrup et al.,1996) and the structure has also been determined, all of which bear evidence that a single unique and biologically active folded product had indeed been produced.

Analytical gelfiltration analysis of rTN proteins.

Analytical gelfiltration of the recombinant tetranectin derivatives TN123, TN3, and TN23 (Fig. 9) were performed on a Superose 12 HR 10/30 column (Pharmacia, Sweden) with a total volume of 25 ml in 100 mM NaCl and 50 mM Tris-HCl pH 8 and a flow rate of 0.2 ml/min. The K_{av} value is defined by, $K_{av} = (Ve-Vo)/(Vc-Vo)$.

The gelfiltration analysis of TN123 and TN23 show that both proteins are exclusively found as trimers in solution (K_{av} values of 0.27 and 0.29, respectively), whereas TN3 appeared monomeric (K_{av}:0.41).

Chemical cross-linking of tetranectin and derivatives

The recombinant tetranectin derivatives TN123, TN3, and TN23, together with the fusion proteins CIIH6FXTN123 and H6FXTN12 or mixtures of these derivatives at 1 mg/ml concentrations in 15 cross-linking buffer (0.1 M Sodium borate, pH 9.1) were incubated with dimethylsuberimidate (DMSI, Sigma). 10 μ l aliquots of protein solution were incubated with 1 μ l aliquots of DMSI stock solution (20 mg/ml in cross-linking buf-20 fer) for 30 minutes at 25°C before addition of 2 μ l quenching buffer (3 M Tris-HCl, pH 9). Subunit exchange between preformed homo-oligomers was induced by subjecting protein mixtures to heat shock treatment. Five μ l aliquots of each protein solution (1 mg/ml stocks) were mixed at 0°C in stan-25 dard polypropylene microcentrifuge tubes, transferred to a water bath at 70°C for the time spans indicated, and then further incubated for 15 minutes at 25°C before reaction with DMSI.

Prior to analysis by SDS-PAGE (12% gels) of the cross-linked products the reaction samples were boiled in the presence of SDS and mercaptoethanol.

Cross-linking analysis of TN123 and the fusion protein CIIH6FXTN123 showed that no detectable subunit exchange between pre-formed homo-oligomers in a mixture of TN123 and CIIH6FXTN123 was found after 16 hours at room temperature 5 (Fig. 10). Subunit exchange could be induced by incubating the protein mixture at 70°C for 15 seconds or longer before cooling to room temperature and addition of DMSI. SDS-PAGE analysis showed the presence of four trimer bands above 95 kDa (corresponding to two homo-trimers and two hetero-tri-10 mers) and three dimer bands (corresponding to two homo-dimers and one hetero-dimer) in the gel between 43 and 55 kDa, in a relative abundance in agreement with random association of monomer subunits into trimers after subunit exchange. It should be noted, that molecular weight markers have only been 15 included on the SDS-PAGE gels for crude calibration and orientation of the gels.

The trimeric organization of tetranectin was further corroborated by cross-linking studies of the proteins H6FXTN12, TN23, and TN3 and mixtures between them (Fig. 11). The tetra-20 nectin derivative TN3, containing only the CRD, could not be cross-linked even at high protein concentrations and did not interfere with the cross-linking of rTN123. Likewise, the derivative TN23, containing exon 2 and the CRD, appeared monomeric after cross-linking and was found not to interfere with trimerisation of TN123 during subunit exchange. Dimeric 25 TN23 molecules found at low abundance in the sample probably reflects contaminating misfolded disulphide bridged dimers. The fusion protein H6FXTN12 formed homo-trimers upon crosslinking and generated hetero-trimers with TN123 after subunit 30 exchange. Because of the difference in size of full length tetranectin (TN123) and H6FXTN12 the possible nine protein bands resulting from chemical cross-linking are: The four trimers [(TN123)₃, (TN123)₂(H6FXTN12), (TN123)(H6FXTN12)₂, and (H6FXTN12)3] at approx. 95 kDa, 50 kDa, 37 kDa, and 20 35 kDa, respectively; the three dimers [(TN123)2, (TN123)(H6FXTN12), and $(H6FXTN12)_2$] at approx. 45 kDa, 30

kDa, and 15 kDa, respectively; and the two monomers TN123 at 23 kDa and H6FXTN12 at 9 kDa.

Taken together, the gel filtration and the cross-linking analysis of the tetranectin derivatives show that tetranectin, like the collectin group of C-type lectins, is a trimeric molecule and that amino acid residues directly shown to be involved in trimerisation of the tetranectin monomer are located in exon 2 of the protein (Val17 - Val49). Furthermore subunit exchange between the trimeric molecules could only be observed after heat denaturation. Amino acid residues Glu1 to Asp16 of tetranectin are critical to chemical cross-linking with DMSI and more important appear to stabilize the trimeric molecule because the cross-linking analysis of the mixture TN123 and TN23 showed no decrease in TN123 formation after heat denaturation and possible subunit 15 exchange (Fig. 11). The stability of the tetranectin trimer was corroborated by a cross-linking analysis with DMSI at different temperatures. Fifteen μ l TN123 at 0.3 mg/ml concentration was pre-incubated 10 min. at either 37°C, 50°C, 60°C, 20 or 70°C before addition of 2 μ l DMSI (20 mg/ml). The reaction was allowed to proceed for 15 min. before reaction was quenched with 5 μ l of 3M Tris-HCl pH 9.1 and the reaction mixtures allowed to cool to room temperature. SDS-PAGE analysis of reduced samples (Fig. 12) showed, that trimers are readily detectable even at 60°C, although a competing pattern 25 of cross-linking specimens increases at increasing temperatures. The appearance of other cross-linking specimens is probably due to the unfolding of the CRD. The stability of the tetranectin trimerising structural element is further analyzed using a designed chimeric protein in Example 3. 30

EXAMPLE 3

Design and construction of the recombinant chimeric protein TRIPB-UB - the tetranectin trimerising structural element and ubiquitin.

5 A plasmid clone, pLCMHF/UB, generously provided by Dr. O. Wiborg harbouring a human ubiquitin cDNA insert (SEQ ID: 16) was used as template and SEQ ID NO: 17 together with SEQ ID NO: 18 were used for amplification in a Polymerase Chain Reaction (PCR) (Saiki et al., 1988) of the nucleotide sequence encoding amino acid residue Ile1 to Gly76 of human ubiquitin(SEQ ID: 19). The amplified DNA fragment was after digestion with the restriction endonucleases BamHI and HindIII ligated into the BamHI and HindIII sites of pTtripb (Example 1) yielding pTtripb-UB (Fig. 13) using standard procedures.

To prepare the chimeric fusion protein H6FXtripb-UB (Fig. 14, SEQ ID NO: 31) the plasmid pTtripb-UB was grown in medium _ scale (4 x 1 litre; 2xTY medium, 5 mM MgSO₄ and 100 μ g ampicillin) in E. coli BL21 cells, as described by Studier et 20 al. (1990). Exponentially growing cultures at 37°C were at OD₆₀₀ 0.8 infected with bacteriophage lambda CE6 at a multiplicity of approximately 5. Cultures were grown at 37°C for another three hours and the cells harvested by centrifugation. Cells were resuspended in 150 ml of 0.5 M NaCl, 10 mM Tris-HCl pH 8, and 1 mM EDTA pH 8. Phenol (100 ml adjusted to pH 8) was added and the mixture sonicated to extract the total protein. Protein was precipitated from the phenol phase by 2.5 volumes of ethanol and centrifugation. The protein pellet was dissolved in a buffer containing 6 M guanidinium chloride, 50 mM Tris-HCl pH 8 and 0.1 M dithioerythriol. Following gel filtration on Sephadex G-25 (Pharmacia, Sweden) into 8 M Urea, 1 M NaCl, 50 mM Tris-HCl pH 8 and 10 mM 2-mercaptoethanol, the crude protein preparation was applied to a Ni²⁺ activated NTA-agarose column for purification (Hochuli et al., 1988) and refolding of the fusion protein H6FXtripb-UB.

Synthesis and charging of the Ni²⁺ activated NTA-agarose matrix is described in Example 2. All buffers for liquid chromatography were degassed prior to use. The fusion protein H6FXtripb-UB was refolded by removing the urea and 2-mercaptoethanol from buffer II in a gradient over 5 column volumes into 50 mM Tris-HCl pH 8 and 0.5 M NaCl. After completion of the refolding procedure the H6FXtripb-UB fusion protein was eluted from the Ni²⁺NTA-agarose columns with a buffer containing 0.5 M NaCl, 50 mM Tris-HCl, 25 mM EDTA pH 8 and gel filtrated on a Sephadex G50 column (Pharmacia) into 0.1 M Sodium borate pH 9 buffer for chemical cross-linking analysis with DMSI.

The cross-linking analysis experiment was designed both to analyze the oligomeric status of the chimeric fusion protein and the thermal stability of the presumed fusion protein trimer as described in Example 2: Samples of 15 μ l H6FXtripb-UB fusion protein, at approximately 1.0 mg/ml concentration, 20 were pre-incubated 10 min. at either 37°C, 50°C, 60°C, or 70°C before addition of 2 μl DMSI (20 mg/ml). The reactions were allowed to proceed for 15 min. before quenching by addition of 5 μ l of 3 M Tris-HCl pH9.1 and the reaction . mixtures were allowed to cool to room temperature. SDS-PAGE analysis of reduced samples (Fig. 12) showed, (1) that the 25 fusion protein H6FXtripb-UB is a trimer in solution (monomer at 17 kDa, dimer at 35 kDa, and trimer at 43 kDa) and (2) that a substantial amount of trimer molecules is present even at 70°C. The appearance of other larger cross-linking products is probably due to cross-linking of trimers via the 30 ubiquitin part of the fusion protein.

EXAMPLE 4

Design and construction of trimerised and hexamerized CEA6 scFv antibodies scFv(CEA6)-TRIPB, TRIPB-scFv(CEA6) and scFv(CEA6)-TRIPB-scFv(CEA6).

A plasmid clone, pUC19MCH/CEA6, generously provided by Dr.
Kevin Pritchard, Cambridge Antibody Technology Ltd., Melbourn, UK, harbouring a nucleotide sequence (SEQ ID: 20)
encoding the CEA6 antibody in single-chain Fv (scFv) format,
followed in sequence by a "myc tag" (which is a general
purification/detection handle), was used as template in
Polymerase Chain Reactions (PCR) (Saiki et al., 1988) in
which the nucleotide sequence encoding the scFv + myc tag was
amplified using the primer pairs (SEQ ID: 21 and SEQ ID: 22)
and (SEQ ID: 21 and SEQ ID: 23) to generate PCR fragments "A"

15 and "B".

PCR fragment "A" was treated with restrictions enzymes BamHI and KpnI and the resulting fragment was inserted into BglII/KpnI cut pTripb (Example 1) to obtain the vector pTH6FXscFv(CEA6)-tripb (Fig. 15) encoding the H6FXscFv(CEA6)-20 TRIPB fusion protein (Fig. 16). PCR fragment "B" was treated with restriction enzymes BamHI and HindIII and the resulting fragment was inserted into BamHI and HindIII cut pTripb (Example 1) to obtain the vector pTH6FXtripb-scFv(CEA6) (Fig. 17) encoding the H6FXTRIPB-scFv(CEA6) fusion protein (Fig.18, SEQ ID NO: 33) using standard procedures.

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To generate the expression vector pTH6FXscFv(CEA6)-tripb-scFv(CEA6) (Fig. 19) encoding the H6FXscFv(CEA6)-TRIPB-scFv(CEA6) fusion protein (Fig. 20, SEQ ID NO: 34) the insert in the vector pTH6FXtripb-scFv(CEA6) was excised using restriction enzymes BamHI and HindIII and inserted into the vector pTH6FXscFv(CEA6)-tripb, which had been treated with restriction enzymes BamHI and HindIII.

To prepare the chimeric fusion proteins H6FXscFv(CEA6)-TRIPB (SEQ ID NO: 32), H6FXTRIPB-scFv(CEA6) (SEQ ID NO: 33) and H6FXscFv(CEA6)-TRIPB-scFv(CEA6) (SEQ ID NO: 34) the plasmids pTH6FXscFv(CEA6)-TRIPB, pTH6FXtripb-scFv(CEA6) and pTH6FXscFv(CEA6)-tripb- scFv(CEA6) were grown in small scale (1 litre; 2xTY medium, 5 mM MgSO4 and 100 μ g ampicillin) in E. coli BL21 cells, as described by Studier et al. (1990). Exponentially growing cultures at 37°C were at OD_{600} 0.8 infected with bacteriophage lambda CE6 at a multiplicity of 10 approximately 5. Cultures were grown at 37°C for another three hours and the cells harvested by centrifugation. Cells were resuspended in 50 ml of 0.5 M NaCl, 50 mM Tris-HCl pH 8, and 1 mM EDTA pH 8. Phenol (50 ml adjusted to pH 8) was added to each and the mixtures were sonicated to extract total protein. After clarification by centrifugation (25 minutes at 10.000 g) crude protein fractions were precipitated from the phenol phases by addition of 2.5 volumes of ethanol and centrifugation. Protein pellets were dissolved in a buffer (15-25 ml) containing 6 M quanidinium chloride, 50 mM 20 Tris-HCl pH 8 and 0.1 M dithioerythriol. Following gel filtration on Sephadex G-25 (Pharmacia, Sweden) into 8 M Urea, 1 M NaCl, 50 mM Tris-HCl pH 8 and 10 mM 2-mercaptoethanol, the crude protein preparations were applied to Ni²⁺ activated NTA-agarose columns (75 ml column volume) for purification 25 (Hochuli et al., 1988). Washing buffer (6 M guanidine-HCl, 50 mM Tris-HCl pH 8 and 10 mM 2-mercaptoethanol) was then flowed through the columns until stable baselines were obtained. Virtually pure fusion proteins could then be eluted by applying a pH gradient to each column (1000 ml gradient in 8 M 30 urea and 10 mM 2-mercaptoethanol obtained by linear (per volume) mixing of solutions containing 50 mM sodium di-hydrogenphosphate (pH 5 buffer) and 50 mM di-sodium hydrogenphosphate (pH 8 buffer).

In preparation for in vitro refolding by the method of Thø35 gersen et al. (WO 94/18227) 20 mg of each purified fusion
protein were mixed in suspensions in refolding "buffer B"
(described below) with aliquots of suspensions of Ni²⁺ acti-

vated NTA-agarose matrix sufficient to generate columns of about 75 ml packed bed volume. Each fusion protein was then subjected to the iterative refolding procedure as described for plasminogen kringle 4 in the Thøgersen et al. patent application (WO 94/18227), except that refolding of the scFv containing fusion proteins was carried out at 10°C using a buffer containing 0.5 M NaCl, 50 mM Tris-HCl pH 8, 2 mM glutathione and 0.2 mM oxidized glutathione as "buffer A" and a buffer containing 8 M urea, 1 M NaCl, 50 mM Tris-HCl pH 8 and 2 mM glutathione as "buffer B".

After completion of the refolding procedure each column was washed with 300 ml buffer containing 0.5 M NaCl and 50 mM Tris-HCl pH 8 to wash away glutathione. The refolded fraction of each protein was then eluted from the NTA-agarose matrix 15 by addition of 20 mM EDTA to the elution buffer. After addition of solid urea to achieve a final concentration of about 8 M to each protein sample and dilution or dialysis to reduce NaCl concentrations to below 5 mM, final purification of each correctly folded fusion protein product was then accomplished 20 by ion exchange chromatography (S-Sepharose, Pharmacia, 1,6 (i.d.) by 90 centimeter column in a buffer containing 8 M urea, 5 mM Tris-HCl (from 1 M stock solution at pH 8) and 25 mM sodium acetate (from 1 M stock solution at pH 5), eluted at 2 ml/min). After dialysis against aqueous buffers (e.g. phosphate buffered saline) each pure and correctly refolded 25 fusion protein was recovered in yields of 2-6 mg per litre of culture grown. Each protein may be shown by analytical gel filtration, chemical cross-linking analysis, by in vitro affinity measurements and by in vivo efficacy to form a 30 stable homotrimeric molecular complex: The oligomeric status of the H6FXtripb-scFv-(CEA6) fusion protein was analyzed by chemical cross-linking analysis with DMSI: In parallel experiments, samples of H6FXtripb-scFv-(CEA6) at 0.34 mg/ml and TN123 at 0.28 mg/ml in 0.1 M Sodium borate were incubated at room temperature with increasing amounts (0 - 40 μ g in 12 μ l in total) of DMSI for 30 min. Reactions were quenched by addition of 5 μ l 3M Tris-HCl pH 9 and the samples analyzed by

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SDS-PAGE under reducing conditions (Fig. 21). Like tetranectin, the H6FXtripb-scFV-(CEA6) fusion protein, of approximately 38 kDa, is hereby shown to be a trimer in solution.

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SEQUENCE LISTING

(1) GENERAL INFORMATION

(i)	APPLICANT
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- (A) NAME: Thøgersen, Hans Christian
- (B) STREET: Ristrupvej 41
- (C) CITY: Mundelstrup
- (D) STATE OR PROVINCE:
- (E) COUNTRY: Denmark
- (F) POSTAL CODE: 8381
- (ii) TITLE OF THE INVENTION: Trimerising module
- (iii) NUMBER OF SEQUENCES: 34
- (iv) COMPUTER-READABLE FORM:
 - (A) MEDIUM TYPE: Diskette
 - (B) COMPUTER: IBM Compatible
 - (C) OPERATING SYSTEM: DOS
 - (D) SOFTWARE: FastSEQ for Windows Version 2.0
- (v) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (2) INFORMATION FOR SEQ ID NO:1:
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 47 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

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47

- (2) INFORMATION FOR SEQ ID NO:2:
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 33 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
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 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single

(D) TOPOLOGY: linear

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 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

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- (2) INFORMATION FOR SEQ ID NO:5:
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 52 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

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 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

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19465DK1.P01/PK/199706 11

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 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

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 - (A) LENGTH: 39 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

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(2) INFORMATION FOR SEQ ID NO:9:

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 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: None
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

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- (2) INFORMATION FOR SEQ ID NO:12:
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 10 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

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19465DK1.P01/PK/199706 11

(2) INFORMATION FOR SEQ ID NO:13:	
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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
CCGAAGCTTA GACCGTCTGC AGGGC	25
(2) INFORMATION FOR SEQ ID NO:14:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 39 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	
GGCGGATCCA TCCAGGGTAG GGTTGTGAAC ACAAAGATG	39
(2) INFORMATION FOR SEQ ID NO:15:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 36 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:	
CCTGGATCCA TCGAGGGTAG GGCCCTGCAG ACGGTC	36
(2) INFORMATION FOR SEQ ID NO:16:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 227 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:	
ATGCAGATCT TTGTGAAGAC CCTCACTGGC AAAACCATCA CCCTTGAGGT CGAGCCCAGT GACACCATTG AGAATGTCAA AGCCAAAATT CAAGACAAGG AGGGTATCCC ACCTGACCGC AGCGTCTGAT ATTTGCCGGC AAACAGCTGG AAGATGGACG TACTTTGTCT GACTACAATA ITCAAAAAGGA GTCTACTCTT CATCTTGTGT TGAGACTTCG TGGTGGT	60 120 180 227
(2) INFORMATION FOR SEO ID NO:17:	

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 27 base pairs(B) TYPE: nucleic acid

(C) STRANDE (D) TOPOLOG	DNESS: single Y: linear		·	
(xi) SEQUENCE	DESCRIPTION:	SEQ ID NO:17:		
TGCTGATCAC AGATCTTT	GT GAAGACC			27
(2) INFORM	MATION FOR SEQ	ID NO:18:		
(A) LENGTH: (B) TYPE: n	CHARACTERISTIC 39 base pairs ucleic acid DNESS: single Y: linear		·	
(xi) SEQUENCE	DESCRIPTION:	SEQ ID NO:18:		
CGCAAGCTTG CATGCTTA	GG ATCCACCACG	AAGTCTCAA		39
(2) INFORM	ATION FOR SEQ	ID NO:19:		
(A) LENGTH: (B) TYPE: a	DNESS: single Y: linear			
(xi) SEQUENCE	DESCRIPTION:	SEQ ID NO:19:		
Met Gln Ile Phe Val	Lys Thr Leu T	hr Gly Lys Thr	Ile Thr Leu 15	Glu
Val Glu Pro Ser Asp 20	Thr Ile Glu A	sn Val Lys Ala		Asp
Lys Glu Gly Ile Pro		ln Arg Leu Ile		Lys
Gln Leu Glu Asp Gly 50	Arg Thr Leu So	er Asp Tyr Asn 60	Ile Gln Lys	Glu
Ser Thr Leu His Leu 65	Val Leu Arg Le 70			
(2) INFORM	ATION FOR SEQ	ID NO:20:		
(A) LENGTH: (B) TYPE: n	CHARACTERISTIC: 786 base pair: ucleic acid DNESS: single Y: linear			

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

CAGGTTCAGC TGCAGCAGTC AGGGGCTGAG GTGAAGAAGC CTGGGTCCTC GGTGAAGGTC 60

TCCTGCAAGG CTTCTGGAGG CACCTTCAGC AACTCTCTA TCAACTGGCT GCGACAGGCC CCCGGACAAG GGCTTGAGTG GATGGAAGT ATCATCCTT CCTTTGGTAC AGCAAACTAC GCTCAGAAGT TCCAGGGCG ACTCACGAT ACCGCGGACG AATCCACGAG CACAGCCTAC ATGGAGCTGA GCAGCCTGAG ATCTACTAC ACCGCGGACG AATCCACGAG CACAGCCTAC ATGGACTGA GCACACTACAC ACCTCTACTA TTACTACATG GACGTCTGGG GCCAGGGGAC AATGGTCACC GTCTCGAGTG GTGGAGGCGG TTCAGGCGGA GGTGGCAGC GCGGTGGCGG ATCGGACATC CACCCTGTCT GCATCTATTG GAGACAGAGT CACCATCACC AGTCTCCTTC CACCCTGTCT GCATCTATTG GAGACAGAGT CACCATCACC GCCCCTAAAC TCCTGATCTA TAAGGCCTCT AGTTTAGCCA GTGGGGCCC ATCAAGGTTC AGCGCAGTG GATCTGGAAC ACAATATAGT AATTATCCGC TCACCTTCG CGGAGGACA GCCTGATGAT TTTGCAACTT ATTACTGCA ACAATATAGT AATTATCCGC TCACCTTCGG CGGAGGACC AGGCCCC AGCAGAGAG GCCAGAGAC CAAAAACTCA TCTCAGAAGA GCAGAGGACC GGGGGCCC AGCACTGAAC GCCGCAGAA ACAATATAGT AATTATCCGC TCACCTTCGG CGGAGGGACC AGGCCCC ACCAACACTCA CCAAAAACTCA TCTCAGAAGA GCAGAGGACC AGGCCCC AGCAGAAACTCA TCTCAGAAGA GCAGAGGACC AGGCCCC ACAAAAACTCA TCTCAGAAGA GCAGAGGACC AGGGGCCC ACAAAAACTCA TCTCAGAAGA GCAGAGAT CAAAAACTCA TCTCAGAAGA GCAGAGACC AGGGGCCC AGGGGCCC AGGGGCCC AGAAAACTCA TCTCAGAAGA GCATCTGAAT GGGGGCCC	120 180 240 300 360 420 480 540 600 660 720 780				
(2) INFORMATION FOR SEQ ID NO:21:					
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 					
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:					
GGTGGATCCC AGGTTCAGCT GCAGC	25				
(2) INFORMATION FOR SEQ ID NO:22:					
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 25 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear					
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:					
GCCGGTACCG GCCCCATTCA GATCC	25				
(2) INFORMATION FOR SEQ ID NO:23:					
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 					
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:					
CCAAGCTTA GGCCCCATTC AGATCC					
(2) INFORMATION FOR SEQ ID NO:24:					
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 228 amino acids					

(B) TYPE: amino acid

- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:

Met Val Arg Ala Asn Lys Arg Asn Glu Ala Leu Arg Ile Glu Ser Ala Leu Leu Asn Lys Ile Ala Met Leu Gly Thr Glu Lys Thr Ala Glu Gly 25 Gly Ser His His His His His Gly Ser Ile Glu Gly Arg Gly Glu Pro Pro Thr Gln Lys Pro Lys Lys Ile Val Asn Ala Lys Lys Asp Val Val Asn Thr Lys Met Phe Glu Glu Leu Lys Ser Arg Leu Asp Thr Leu Ala Gln Glu Val Ala Leu Leu Lys Glu Gln Gln Ala Leu Gln Thr Val 90 85 Cys Leu Lys Gly Thr Lys Val His Met Lys Cys Phe Leu Ala Phe Thr 105 Gln Thr Lys Thr Phe His Glu Ala Ser Glu Asp Cys Ile Ser Arg Gly 120 Gly Thr Leu Ser Thr Pro Gln Thr Gly Ser Glu Asn Asp Ala Leu Tyr 135 Glu Tyr Leu Arg Gln Ser Val Gly Asn Glu Ala Glu Ile Trp Leu Gly 150 155 Leu Asn Asp Met Ala Ala Glu Gly Thr Trp Val Asp Met Thr Gly Ala 165 170 Arg Ile Ala Tyr Lys Asn Trp Glu Thr Glu Ile Thr Ala Gln Pro Asp 185 Gly Gly Lys Thr Glu Asn Cys Ala Val Leu Ser Gly Ala Ala Asn Gly 200 205 Lys Trp Phe Asp Lys Arg Cys Arg Asp Gln Leu Pro Tyr Ile Cys Gln 215 Phe Gly Ile Val 225

(2) INFORMATION FOR SEQ ID NO:25:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 197 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:

 Met Gly Ser His His His His His His His Gly Ser Ile Glu Gly Arg Gly

 1
 5
 10
 15

 Glu Pro Pro Thr Gln Lys Pro Lys Lys Ile Val Asn Ala Lys Lys Asp
 20
 25
 30

 Val Val Asn Thr Lys Met Phe Glu Glu Leu Lys Ser Arg Leu Asp Thr
 35
 40
 45

 Leu Ala Gln Glu Val Ala Leu Leu Lys Glu Gln Gln Ala Leu Gln Thr

Val Cys Leu Lys Gly Thr Lys Val His Met Lys Cys Phe Leu Ala Phe Thr Gln Thr Lys Thr Phe His Glu Ala Ser Glu Asp Cys Ile Ser Arg 85 90 Gly Gly Thr Leu Ser Thr Pro Gln Thr Gly Ser Glu Asn Asp Ala Leu 100 105 Tyr Glu Tyr Leu Arg Gln Ser Val Gly Asn Glu Ala Glu Ile Trp Leu 120 Gly Leu Asn Asp Met Ala Ala Glu Gly Thr Trp Val Asp Met Thr Gly 135 140 Ala Arg Ile Ala Tyr Lys Asn Trp Glu Thr Glu Ile Thr Ala Gln Pro 150 155 Asp Gly Gly Lys Thr Glu Asn Cys Ala Val Leu Ser Gly Ala Ala Asn 165 170 Gly Lys Trp Phe Asp Lys Arg Cys Arg Asp Gln Leu Pro Tyr Ile Cys 185 Gln Phe Gly Ile Val 195

(2) INFORMATION FOR SEQ ID NO:26:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 65 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:

 Met
 Gly
 Ser
 His
 His
 His
 His
 His
 His
 Gly
 Ser
 Ile
 Glu
 Gly
 Arg
 Gly
 Ile
 I

- (2) INFORMATION FOR SEQ ID NO:27:
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 180 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:

Met Gly Ser His His His His His Gly Ser Ile Gln Gly Arg Val

1 5 10 15

Val Asn Thr Lys Met Phe Glu Glu Leu Lys Ser Arg Leu Asp Thr Leu
20 25 30

Ala Gln Glu Val Ala Leu Leu Lys Glu Gln Gln Ala Leu Gln Thr Val Cys Leu Lys Gly Thr Lys Val His Met Lys Cys Phe Leu Ala Phe Thr Gln Thr Lys Thr Phe His Glu Ala Ser Glu Asp Cys Ile Ser Arg Gly 70 75 Gly Thr Leu Ser Thr Pro Gln Thr Gly Ser Glu Asn Asp Ala Leu Tyr 85 90 Glu Tyr Leu Arg Gln Ser Val Gly Asn Glu Ala Glu Ile Trp Leu Gly 100 105 Leu Asn Asp Met Ala Ala Glu Gly Thr Trp Val Asp Met Thr Gly Ala 120 Arg Ile Ala Tyr Lys Asn Trp Glu Thr Glu Ile Thr Ala Gln Pro Asp 135 140 Gly Gly Lys Thr Glu Asn Cys Ala Val Leu Ser Gly Ala Ala Asn Gly 155 Lys Trp Phe Asp Lys Arg Cys Arg Asp Gln Leu Pro Tyr Ile Cys Gln 165 170 Phe Gly Ile Val 180

(2) INFORMATION FOR SEQ ID NO:28:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 73 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:

 Met Gly Ser His His His His His His His His Gly Ser Ile Gln Gly Arg Ser

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 10
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 Pro Gly Thr Glu Pro Pro Thr Gln Lys Pro Lys Lys Ile Val Asn Ala 20
 25
 30

 Lys Lys Asp Val Val Asn Thr Lys Met Phe Glu Glu Leu Lys Ser Arg 35
 40
 45

 Leu Asp Thr Leu Ala Gln Glu Val Ala Leu Leu Lys Glu Glu Gln Gln Ala 50
 55
 60

 Leu Gln Thr Val Ser Leu Lys Gly Ser 65
 70

(2) INFORMATION FOR SEQ ID NO:29:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 69 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:

Met Gly Ser His His His His His Gly Ser Ile Gln Gly Arg Ser

1 10 15

 Pro
 Gly
 Gly
 Pro
 Pro
 Pro
 Lys
 Pro
 Lys
 L

(2) INFORMATION FOR SEQ ID NO:30:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 152 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:

Met Gly Ser His His His His His Gly Ser Ile Glu Gly Arg Ala Leu Gln Thr Val Cys Leu Lys Gly Thr Lys Val His Met Lys Cys Phe 25 Leu Ala Phe Thr Gln Thr Lys Thr Phe His Glu Ala Ser Glu Asp Cys Ile Ser Arg Gly Gly Thr Leu Ser Thr Pro Gln Thr Gly Ser Glu Asn Asp Ala Leu Tyr Glu Tyr Leu Arg Gln Ser Val Gly Asn Glu Ala Glu 75 Ile Trp Leu Gly Leu Asn Asp Met Ala Ala Glu Gly Thr Trp Val Asp 90 Met Thr Gly Ala Arg Ile Ala Tyr Lys Asn Trp Glu Thr Glu Ile Thr 105 Ala Gln Pro Asp Gly Gly Lys Thr Glu Asn Cys Ala Val Leu Ser Gly 120 Ala Ala Asn Gly Lys Trp Phe Asp Lys Arg Cys Arg Asp Gln Leu Pro 135 Tyr Ile Cys Gln Phe Gly Ile Val 145 150

- (2) INFORMATION FOR SEQ ID NO:31:
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 145 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:

Met Gly Ser His His His His His Gly Ser Ile Gln Gly Arg Ser 1 5 10 15

Pro Gly Thr Glu Pro Pro Thr Gln Lys Pro Lys Lys Ile Val Asn Ala 20 25 30

Lys Lys Asp Val Val Asn Thr Lys Met Phe Glu Glu Leu Lys Ser Arg 40 Leu Asp Thr Leu Ala Gln Glu Val Ala Leu Leu Lys Glu Gln Gln Ala 55 Leu Gln Thr Gly Ser Gln Ile Phe Val Lys Thr Leu Thr Gly Lys Thr 75 70 Ile Thr Leu Glu Val Glu Pro Ser Asp Thr Ile Glu Asn Val Lys Ala Lys Ile Gln Asp Lys Glu Gly Ile Pro Pro Asp Gln Gln Arg Leu Ile 105 Phe Ala Gly Lys Gln Leu Glu Asp Gly Arg Thr Leu Ser Asp Tyr Asn 120 125 Ile Gln Lys Glu Ser Thr Leu His Leu Val Leu Arg Leu Arg Gly Gly 135 140 Ser 145

(2) INFORMATION FOR SEQ ID NO:32:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 330 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:

Met Gly Ser His His His His His Gly Ser Ile Gln Gly Arg Ser Gln Val Gln Leu Gln Gln Ser Gly Ala Glu Val Lys Lys Pro Gly Ser Ser Val Lys Val Ser Cys Lys Ala Ser Gly Gly Thr Phe Ser Asn Ser Pro Ile Asn Trp Leu Arg Gln Ala Pro Gly Gln Gly Leu Glu Trp Met 55 Gly Ser Ile Ile Pro Ser Phe Gly Thr Ala Asn Tyr Ala Gln Lys Phe 70 Gln Gly Arg Leu Thr Ile Thr Ala Asp Glu Ser Thr Ser Thr Ala Tyr Met Glu Leu Ser Ser Leu Arg Ser Glu Asp Thr Ala Val Tyr Tyr Cys Ala Gly Arg Ser His Asn Tyr Glu Leu Tyr Tyr Tyr Met Asp Val 120 125 Trp Gly Gln Gly Thr Met Val Thr Val Ser Ser Gly Gly Gly Ser 135 Gly Gly Gly Ser Gly Gly Gly Ser Asp Ile Gln Met Thr Gln 150 155 Ser Pro Ser Thr Leu Ser Ala Ser Ile Gly Asp Arg Val Thr Ile Thr 165 170 Cys Arg Ala Ser Glu Gly Ile Tyr His Trp Leu Ala Trp Tyr Gln Gln 180 185 190 Lys Pro Gly Lys Ala Pro Lys Leu Leu Ile Tyr Lys Ala Ser Ser Leu 200 205 Ala Ser Gly Ala Pro Ser Arg Phe Ser Gly Ser Gly Ser Gly Thr Asp 215 220 210

Phe Thr Leu Thr Ile Ser Ser Leu Gln Pro Asp Asp Phe Ala Thr Tyr 235 Tyr Cys Gln Gln Tyr Ser Asn Tyr Pro Leu Thr Phe Gly Gly Gly Thr 245 250 Lys Leu Glu Ile Lys Arg Ala Ala Glu Gln Lys Leu Ile Ser Glu 265 Glu Asp Leu Asn Gly Ala Gly Thr Glu Pro Pro Thr Gln Lys Pro Lys 280 285 Lys Ile Val Asn Ala Lys Lys Asp Val Val Asn Thr Lys Met Phe Glu 295 Glu Leu Lys Ser Arg Leu Asp Thr Leu Ala Gln Glu Val Ala Leu Leu 310 315 Lys Glu Gln Gln Ala Leu Gln Thr Gly Ser 325

(2) INFORMATION FOR SEQ ID NO:33:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 331 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:

Met Gly Ser His His His His His Gly Ser Ile Gln Gly Arg Ser Pro Gly Thr Glu Pro Pro Thr Gln Lys Pro Lys Lys Ile Val Asn Ala 25 Lys Lys Asp Val Val Asn Thr Lys Met Phe Glu Glu Leu Lys Ser Arg 40 Leu Asp Thr Leu Ala Gln Glu Val Ala Leu Leu Lys Glu Gln Gln Ala Leu Gln Thr Gly Ser Gln Val Gln Leu Gln Gln Ser Gly Ala Glu Val Lys Lys Pro Gly Ser Ser Val Lys Val Ser Cys Lys Ala Ser Gly Gly Thr Phe Ser Asn Ser Pro Ile Asn Trp Leu Arg Gln Ala Pro Gly Gln 105 Gly Leu Glu Trp Met Gly Ser Ile Ile Pro Ser Phe Gly Thr Ala Asn 120 125 Tyr Ala Gln Lys Phe Gln Gly Arg Leu Thr Ile Thr Ala Asp Glu Ser 135 140 Thr Ser Thr Ala Tyr Met Glu Leu Ser Ser Leu Arg Ser Glu Asp Thr 150 155 Ala Val Tyr Tyr Cys Ala Gly Arg Ser His Asn Tyr Glu Leu Tyr Tyr 165 170 Tyr Tyr Met Asp Val Trp Gly Gln Gly Thr Met Val Thr Val Ser Ser 180 185 Gly Gly Gly Ser Gly Gly Gly Gly Ser Gly Gly Gly Ser Asp 200 Ile Gln Met Thr Gln Ser Pro Ser Thr Leu Ser Ala Ser Ile Gly Asp 215 220 Arg Val Thr Ile Thr Cys Arg Ala Ser Glu Gly Ile Tyr His Trp Leu 230 235

(2) INFORMATION FOR SEQ ID NO:34:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 592 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:

Met Gly Ser His His His His His Gly Ser Ile Gln Gly Arg Ser Gln Val Gln Leu Gln Gln Ser Gly Ala Glu Val Lys Lys Pro Gly Ser 25 Ser Val Lys Val Ser Cys Lys Ala Ser Gly Gly Thr Phe Ser Asn Ser Pro Ile Asn Trp Leu Arg Gln Ala Pro Gly Gln Gly Leu Glu Trp Met 55 Gly Ser Ile Ile Pro Ser Phe Gly Thr Ala Asn Tyr Ala Gln Lys Phe 70 75 Gln Gly Arg Leu Thr Ile Thr Ala Asp Glu Ser Thr Ser Thr Ala Tyr 90 Met Glu Leu Ser Ser Leu Arg Ser Glu Asp Thr Ala Val Tyr Tyr Cys 105 Ala Gly Arg Ser His Asn Tyr Glu Leu Tyr Tyr Tyr Tyr Met Asp Val 120 125 Trp Gly Gln Gly Thr Met Val Thr Val Ser Ser Gly Gly Gly Ser 135 Gly Gly Gly Ser Gly Gly Gly Ser Asp Ile Gln Met Thr Gln 150 155 Ser Pro Ser Thr Leu Ser Ala Ser Ile Gly Asp Arg Val Thr Ile Thr 165 170 Cys Arg Ala Ser Glu Gly Ile Tyr His Trp Leu Ala Trp Tyr Gln Gln 185 190 Lys Pro Gly Lys Ala Pro Lys Leu Leu Ile Tyr Lys Ala Ser Ser Leu 200 Ala Ser Gly Ala Pro Ser Arg Phe Ser Gly Ser Gly Ser Gly Thr Asp 215 220 Phe Thr Leu Thr Ile Ser Ser Leu Gln Pro Asp Asp Phe Ala Thr Tyr 230 235 Tyr Cys Gln Gln Tyr Ser Asn Tyr Pro Leu Thr Phe Gly Gly Gly Thr 250

Lys Leu Glu Ile Lys Arg Ala Ala Glu Gln Lys Leu Ile Ser Glu Glu Asp Leu Asn Gly Ala Gly Thr Glu Pro Pro Thr Gln Lys Pro Lys Lys Ile Val Asn Ala Lys Lys Asp Val Val Asn Thr Lys Met Phe Glu Glu Leu Lys Ser Arg Leu Asp Thr Leu Ala Gln Glu Val Ala Leu Leu Lys Glu Gln Gln Ala Leu Gln Thr Gly Ser Gln Val Gln Leu Gln Gln Ser Gly Ala Glu Val Lys Lys Pro Gly Ser Ser Val Lys Val Ser Cys Lys Ala Ser Gly Gly Thr Phe Ser Asn Ser Pro Ile Asn Trp Leu Arg Gln Ala Pro Gly Gln Gly Leu Glu Trp Met Gly Ser Ile Ile Pro Ser Phe Gly Thr Ala Asn Tyr Ala Gln Lys Phe Gln Gly Arg Leu Thr Ile Thr Ala Asp Glu Ser Thr Ser Thr Ala Tyr Met Glu Leu Ser Ser Leu Arg Ser Glu Asp Thr Ala Val Tyr Tyr Cys Ala Gly Arg Ser His Asn Tyr Glu Leu Tyr Tyr Tyr Met Asp Val Trp Gly Gln Gly Thr Met Val Thr Val Ser Ser Gly Gly Gly Gly Ser Gly Gly Gly Ser Gly Gly Gly Ser Asp Ile Gln Met Thr Gln Ser Pro Ser Thr Leu Ser Ala Ser Ile Gly Asp Arg Val Thr Ile Thr Cys Arg Ala Ser Glu Gly Ile Tyr His Trp Leu Ala Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Lys Leu Leu Ile Tyr Lys Ala Ser Ser Leu Ala Ser Gly Ala Pro Ser Arg Phe Ser Gly Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Ser Ser Leu Gln Pro Asp Asp Phe Ala Thr Tyr Tyr Cys Gln Gln Tyr Ser Asn Tyr Pro Leu Thr Phe Gly Gly Gly Thr Lys Leu Glu Ile Lys Arg Ala Ala Ala Glu Gln Lys Leu Ile Ser Glu Glu Asp Leu Asn Gly Ala

CLAIMS

- A monomer polypeptide construct comprising at least one tetranectin trimerising structural element (TTSE) which is covalently linked to at least one heterologous moiety, said
 TTSE being capable of forming a stable complex with two other TTSEs, with the proviso that the heterologous moiety is different from any of the fusion proteins CIIH6FXTN123, H6FXTN123, H6FXTN123, the sequences of which are shown in SEQ ID NOS: 24-27.
- 2. A monomer polypeptide construct comprising at least one tetranectin trimerising structural element (TTSE) which is covalently linked to at least one heterologous moiety, said TTSE being capable of forming a stable complex with two other TTSEs, said at least one heterologous moiety being one which does not exclusively facilitate expression and/or purification of the monomer polypeptide construct.
 - 3. A monomer polypeptide construct according to claim 1 or 2, wherein the stable complex includes a triple alpha helical coiled coil.
- 4. A monomer polypeptide construct according to any of the preceding claims, wherein the heterologous moiety is selected from the group consisting of a ligand binding structure; a toxin; a detectable label; an in situ activatable substance; an enzyme; a radioactive moiety; a cytokine; a non-protein-aceous polymer such as a polymeric alkaloid, a polyalcohol, a polysaccharide, a lipid and a polyamine; a photo cross-linking agent; and a group facilitating conjugation of the monomer polypeptide construct to a target.
- 5. A monomer polypeptide construct according to any of the preceding claims, which comprises 2 TTSEs which are covalently linked by a spacer moiety which allows both of the 2 TTSEs to take part in complex formation with a third TTSE not being part of the monomer polypeptide construct.

- 6. A monomer polypeptide construct according to claim 5, wherein the spacer moiety has a length and a conformation which favours complex formation involving both of the two TTSEs which are covalently linked by the spacer moiety.
- 7. A monomer polypeptide construct according to claim 5 or 6, wherein the spacer moiety is a polypeptide fragment.
 - 8. A monomer polypeptide construct according to any of claims 1-4, which comprises one single TTSE.
- 9. A monomer polypeptide construct according to any of the preceding claims, wherein the TTSE is derived from human tetranectin, murine tetranectin, C-type lectin of bovine cartilage, or C-type lectin of shark cartilage.
- 10. A monomer polypeptide construct according to claim 9, wherein the TTSE comprises a polypeptide sequence which has at least 68% sequence identity with the consensus sequence shown in Fig. 2.
 - 11. A monomer polypeptide construct according to claim 10, wherein the sequence identity with the consensus sequence is at least 75%, such as at least 81%, at least 87%, or at least 92%.
- 12. A monomer polypeptide construct according to any of claims 9-11, wherein at least one amino acid residue selected from the group consisting of amino acid residue nos. 6, 21, 22, 24, 25, 27, 28, 31, 32, 35, 39, 41, 42, is/are substituted by any non-helix breaking amino acid residue, the amino acid residue numbering referring to amino acid residues in
- acid residue numbering referring to amino acid residues in SEQ ID NO: 7.
- 13. A monomer polypeptide construct according to any of the preceding claims, wherein the at least one TTSE comprises a repeated heptad having the formula a-b-c-d-e-f-g (N to C),

wherein residues a and d generally are hydrophobic amino acids.

- 14. A monomer polypeptide construct according to claim 13, wherein heptad is repeated 3 times and wherein the last5 occurrence of the heptad has a glutamine residue corresponding to residues a and d.
- 15. A monomer polypeptide construct according to any of the preceding claims, wherein the heterologous moiety is covalently linked to the TTSE by via a peptide bond to the N- or C-terminus of the TTSE peptide chain, via a peptide bond to a side chain in the TTSE or via a bond to a cysteine residue.
 - 16. A monomer polypeptide construct according to any of the preceding claims which lacks any free amino and/or carboxy groups.
- 15 17. A monomer polypeptide construct according to any of the preceding claims which lacks substantial of tetranectin which is encoded by exon 3 and/or lacks substantial part of tetranectin which is encoded by exon 1.
- 18. A monomer polypeptide construct according to any of the preceding claims comprising two heterologous moieties which are linked via peptide bonds to the N- and C-terminus, respectively.
- 19. A monomer polypeptide construct according to any of the preceding claims which is constructed so as to disfavour 25 formation of complexes between identical TTSEs.
 - 20. An oligomer which is comprised of two monomer polypeptide constructs according to any of claims 1-19, and which comprises at least three TTSE's, or which is comprised of three monomer polypeptide constructs according to any of claims 1-4 or 8-19.

- 23. An oligomer according to claim 22, wherein the at least one heterologous moiety which is positioned N-terminally to a TTSE and the at least one heterologous moiety which is positioned C-terminally to a TTSE are part of the same monomeric polypeptide construct.
- 24. An oligomer according to claim 22, wherein the at least one heterologous moiety which is positioned N-terminally to a TTSE and the at least one heterologous moiety which is positioned C-terminally to a TTSE are part of two separate monomeric polypeptide constructs.
 - 25. An oligomer according to any of claims 20-24, wherein each monomer polypeptide construct is designed so as to disfavour formation of trimers including two monomer polypeptide constructs having identical TTSEs.
 - 26. A method of preparing a monomer polypeptide construct according to any of claims 1-19, the method comprising
- isolating the monomer polypeptide construct from a culture comprising a host cell which carries and expresses a nucleic acid fragment which encodes the monomer polypep-25 tide construct,
 - synthesizing, by means of chemical peptide synthesis, the monomer polypeptide construct and subsequently isolating the monomer polypeptide construct from the reaction mixture,

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- preparing a TTSE in a culture comprising a host cell which carries and expresses a nucleic acid fragment which encodes the TTSE, subsequently linking covalently at least one heterologous moiety to the TTSE, and thereafter isolating the resulting monomer polypeptide construct, or
- synthesizing, by means of chemical peptide synthesis, a TTSE, subsequently linking covalently at least one heterologous moiety to the TTSE, and thereafter the isolating the resulting monomer polypeptide construct from the reaction mixture,

and optionally subjecting the monomer polypeptide construct to further processing.

- 27. A method for preparing a dimeric oligomer according to claim 20 which comprises
- of claims 1-19 which includes two TTSEs (construct 1) with a monomer polypeptide construct according to any of claims 1-4 or 8-19 which includes only one TTSE (construct 2),
- 20 effecting the two TTSE's of construct 1 to complex with the TTSE of construct 2, and
 - isolating the resulting dimer and optionally subjecting the dimer to further processing.
- 28. A method for preparing a trimeric oligomer according to claim 20 which comprises
 - admixing three monomer polypeptide constructs according to any of claims 1-19 with each other,
 - effecting complex formation between one TTSE of each monomer polypeptide construct, and

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- isolating the resulting trimer and optionally subjecting the trimeric oligomer to further processing.

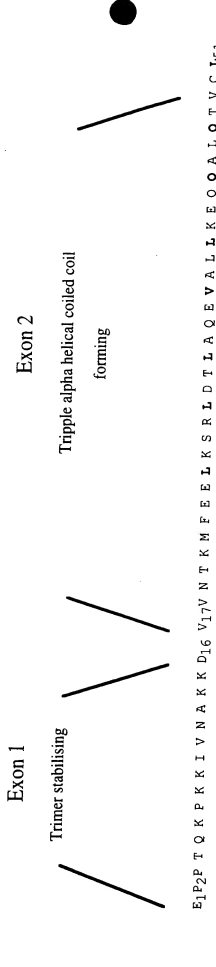
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29. A kit comprising

- a first package comprising at least one container means, 5 each at least one container means containing a monomer polypeptide construct according to any of claims 1-19,
- a second package comprising at least one container means, each at least one container means in the second package containing a monomer polypeptide construct according to any of claims 1-19, the second package being different from the first package with respect to choice and/or number of monomer polypeptide constructs included therein, and optionally
- a third package comprising at least one container means,
 each at least one container means in the third package
 containing a monomer polypeptide construct according to
 any of claims 1-19, the second package being different
 from the first and second packages with respect to choice
 and/or number of monomer polypeptide constructs included
 therein.
 - 30. A kit according to claim 29, wherein the at least one container means in each package contains mutually distinct monomer polypeptide constructs.
- 31. A kit according to claim 29 or 30, wherein all container means comprised in the kit comprises mutually distinct polypeptide constructs.
- 32. A nucleic acid fragment in isolated form which encodes a TTSE as defined in any of claims 1-19 or which encodes the polypeptide part of a monomer polypeptide construct according to any of claims 1-19, with the proviso that the nucleic acid fragment is different from one that encodes native members of

the tetranectin family, and that the nucleic acid fragment is different from one that encodes any of the fusion proteins CIIH6FXTN123, H6FXTN123, H6FXTN12, H6FXTN23, the sequences of which are shown in SEQ ID NOs: 24-27.

- 5 33. A replicable vector which comprises a nucleic acid fragment according to claim 32.
 - 34. A transformed host cell, which comprises a nucleic acid fragment according to claim 32 or a replicable vector according to claim 32.



TLAQEVALLKEQQALQTVCL51

R L D

E L K

Fig. 1

× × α × J đ Ы 1 ч Ç b Ö C Ö > 44 > > > Н 드 Φ H ₽ Ø ש O! Q П Ц U Ц П Ω ď ď ø A O² O Ø ๙ Ø Ø Σ g × Σ ഥ 团 ഠ 山 × **1** ⊠ Φ × × Ч で Н П Ø Ö Ц П ഗ Д Ø ø z Z > ಗ > > > 団 b ſΞÌ 团 ſΞÌ Õ Ø 召 ĸ Φ ď ø 3 3 Н Ы ヷ Н Н O H > × × Д Д Ω ſτĴ Ω ಥ ı Σ > Н 吆 Ø р α 闰 S z z \vdash ĸ K × × **Γ** Ы Н Ы Щ 口 Ω 闰 闰 G О ഥ ഥ Д × Σ Σ × Q ¥ × 闰 × H ഗ × Ö Z വ \gt Ŋ ۸ ۸ □ ĸ S X œ Bovine cart. protein Shark cart. protein Murine tetranectin Human tetranectin Position

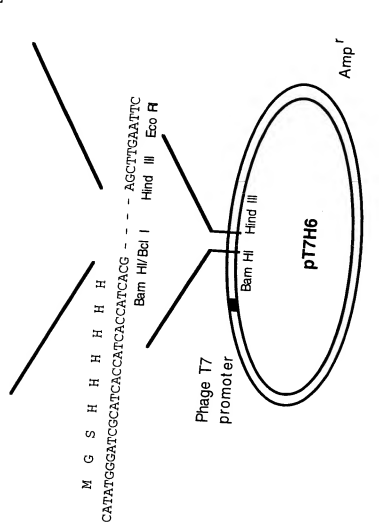
ပ > ⊱ Ø Ц ď Ø 山 × H E Н ЬŻ Ч Consensus

1

Fig. 2

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A L Q T G S * - GCCCTGCAGACGGGATCCTAA Bam H G S I Q G R S P G T E P P T GATCAATCCAGGAAGATCTCCTGGTACCGAGCCACCAACC Kpn ≡ tripb



Ŋ

Fig. 3

H6FXtripa fusion protein

1	M	G	S	Н	Н	H	Н	Н	Н	G	S	I	Q	G	R	S	Р	G	T	E	P	P	T	Q	K	P	K	K	I	V	30
31	N	A	K	K	D	V	V	N	Т	K	M	F	E	E	L	K	S	R	L	D	Т	L	A	Q	E	V	Α	L	L	K	60
61	E	Q	Q	Α	L	Q	Т	v	S	L	K	G	s	*																	73

H6FXtripB fusion protein

1	М	G	S	Н	Н	Н	Н	Н	Η	G	S	Ι	Q	G	R	S	Ρ	G	Т	Ε	Р	P	Т	Q	K	Ρ	K	K	Ι	V	30
31	N	A	K	K	D	V	V	N	Т	K	M	F	E	E	L	K	S	R	L	D	Т	L	A	Q	E	V	A	L	L	K	60
61	E	Q	Q	A	L	Q	т	G	s	*																					69

Fig. 4

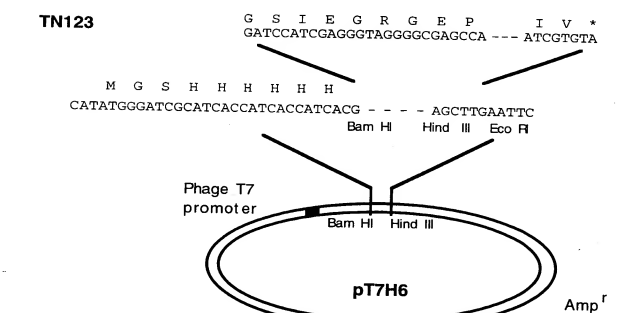


Fig. 5

CIIH6FXTN123 fusion protein

M V R A N K R N E A L R I E S A L L N K I A M L G T E K T A 30 EGGSHНННННСЅІЕСРСРТОКРККІ V N A 31 60 K K D V V N T K M F E E L K S R L D T L A Q E V A L L K E Q 61 90 Q A L Q T V C L K G T K V H M K C F L A F T Q T K T F H E A 120 91 SEDCISRGGTLSTPQTGSENDALYEYLRQS150 121 V G N E A E I W L G L N D M A A E G T W V D M T G A R I A Y 180 151 KNWETEITAQPDGGKTENCAVLSGAANGKW 210 181 F D K R C R D Q L P Y I C Q F G I V * 211 228

H6FXTN123 fusion protein

TN12 G S I E G R G E P P Q T V *
GATCCATCGAGGGTAGGGGGGGGCGAGCCACCA ---- CAGACGGTCTA

TN3

TN23 G S I Q G R V V N T G I V * GATCCATCCAGGGTAGGGTTGTGAACACA ---- GGGATCGTGTA

G S I E G R A L Q G I V *
GATCCATCGAGGGTAGGGCCCTGCAG ---- GGGATCGTGTA

M G S H H H H H H

CATATGGGATCGCATCACCATCACCATCACG - - - AGCTTGAATTC

Bam HI Hind III Eco FI

Phage T7

promoter

Bam HI Hind III

Amp r

Fig. 7

H6FXTN12 fusion protein

1 MGSHHHHHHGSIEGRGEPPTQKPKKIVNAK 30
31 KDVVNTKMFEELKSRLDTLAQEVALLKEQQ 60
61 ALQTV*

H6FXTN23 fusion protein

1 M G S H H H H H H G S I Q G R V V N T K M F E E L K S R L D 30
31 T L A Q E V A L L K E Q Q A L Q T V C L K G T K V H M K C F 60
61 L A F T Q T K T F H E A S E D C I S R G G T L S T P Q T G S 90
91 E N D A L Y E Y L R Q S V G N E A E I W L G L N D M A A E G 120
121 T W V D M T G A R I A Y K N W E T E I T A Q P D G G K T E N 150
151 C A V L S G A A N G K W F D K R C R D Q L P Y I C Q F G I V 180
181 *

H6FXTN3 fusion protein

1 M G S H H H H H H G S I E G R A L Q T V C L K G T K V H M K 30
31 C F L A F T Q T K T F H E A S E D C I S R G G T L S T P Q T 60
61 G S E N D A L Y E Y L R Q S V G N E A E I W L G L N D M A A 90
91 E G T W V D M T G A R I A Y K N W E T E I T A Q P D G G K T 120
121 E N C A V L S G A A N G K W F D K R C R D Q L P Y I C Q F G 150
151 I V *

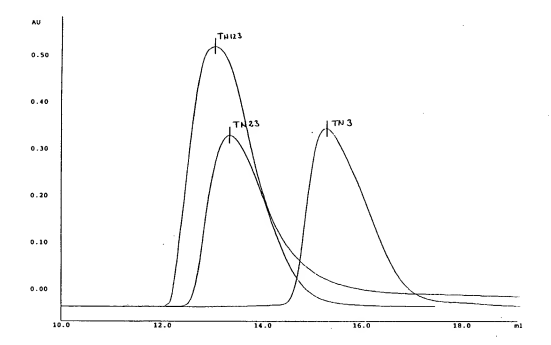


Fig. 9

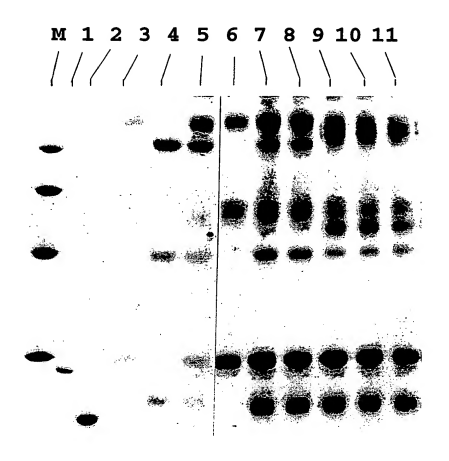


Fig. 10

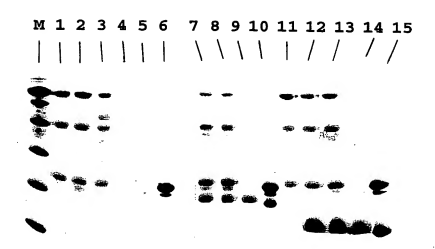


Fig. 11

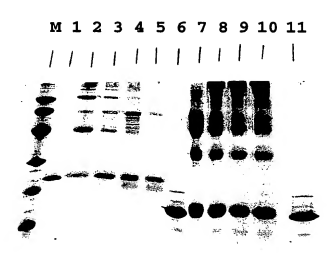


Fig. 12

NB

Fig. 13

Amp

pT7H6

H6FXtripb-UB fusion protein

1 MGSHHHHHHGSIQGRSPGTEPPTQKPKIV 30
31 NAKKDVNNTKMFEELKKSRLDTLAQEVALLK60
61 EQQALQTGSQIFVKTLTGKTITLEVEPSDT90
91 IENVKAKIQKEGIPPDQQRLIFAGKQLED120

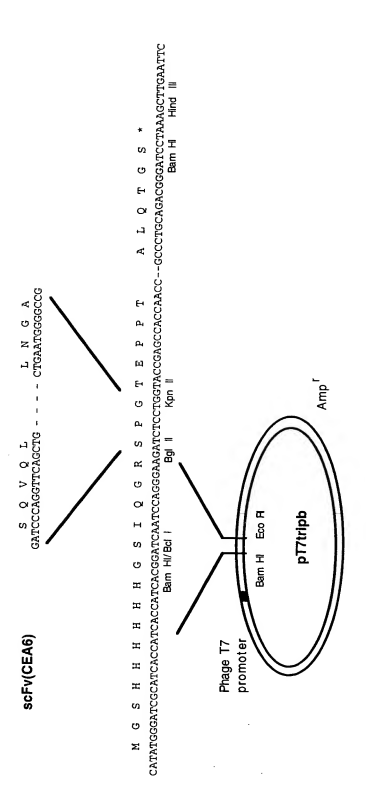


Fig. 15

H6FXscFv(CEA6)-tripb fusion protein

Fig. 16

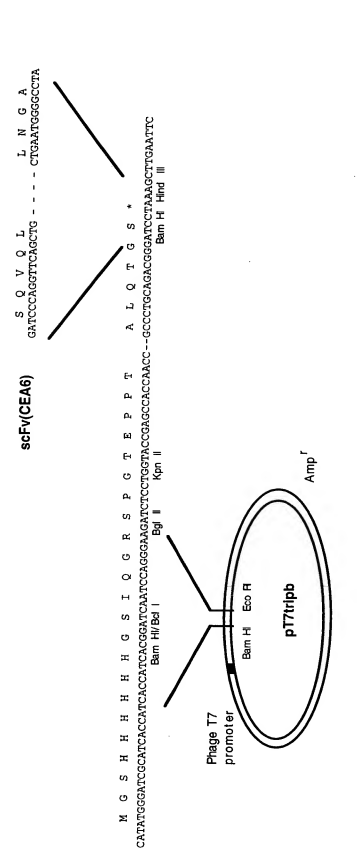


Fig. 17

H6FXtripb-scFv(CEA6) fusion protein

Fig. 18

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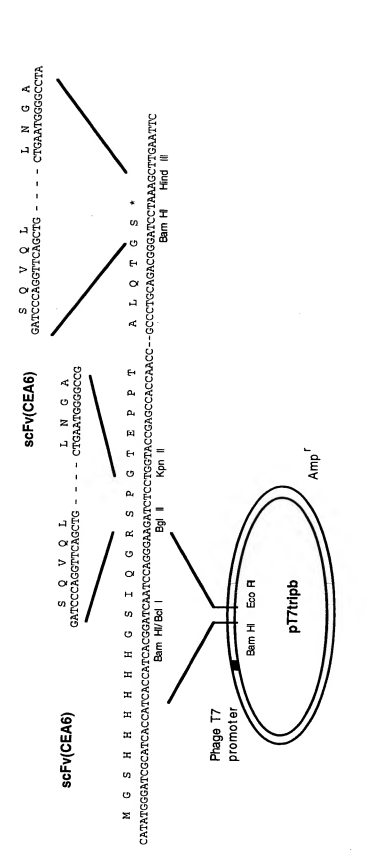


Fig. 19

H6FXscFv(CEA6)-tripb-scFv(CEA6) fusion protein

1 M G S H H H H H H G S I Q G R S Q V Q L Q Q S G A E V K K P 31 G S S V K V S C K A S G G T F S N S P I N W L R Q A P G Q G 60 61 L E W M G S I I P S F G T A N Y A Q K F Q G R L T I T A D E 91 S T S T A Y M E L S S L R S E D T A V Y Y C A G R S H N Y E 120 121 L Y Y Y M D V W G Q G T M V T V S S G G G S G G G S G 150 151 G G G S D I Q M T Q S P S T L S A S I G D R V T I T C R A S 180 181 E G I Y H W L A W Y Q Q K P G K A P K L L I Y K A S S L A S 210 211 GAPSRFSGSGSGTDFTLTISSLQPDDFATY 240 241 Y C Q Q Y S N Y P L T F G G G T K L E I K R A A A E Q K L I 270 271 SEEDLNGAGTEPPTQKPKKIVNAKKDVVNT 300 301 KMFEELKSRLDTLAQEVALLKEQQALQTGS330 331 Q V Q L Q Q S G A E V K K P G S S V K V S C K A S G G T F S 360 361 N S P I N W L R Q A P G Q G L E W M G S I I P S F G T A N Y 390 391 A Q K F Q G R L T I T A D E S T S T A Y M E L S S L R S E D 420 421 TAVYYCAGRSHNYELYYYYMDVWGQGTMVT450 451 V S S G G G G G G G G G G G G G D I Q M T Q S P S T L S 480 481 A S I G D R V T I T C R A S E G I Y H W L A W Y Q Q K P G K 510 511 APKLLIYKASSLASGAPSRFSGSGSGTDFT540 541 LTISSLQPDDFATYYCQQYSNYPLTFGGGT 570 571 K L E I K R A A A E Q K L I S E E D L N G A * 592

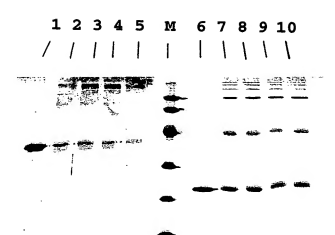


Fig. 21